





CARBOHYDRATE METABOLISM

CORRELATION OF PHYSIOLOGICAL, BIOCHEMICAL AND CLINICAL ASPECTS

By

SAMUEL SOSKIN, M D

Director of the Research Institute Afschool Reese Ho Medical Director, Michael Reese Hospital Professonal Lectures in Physiology University of C.

AND

RACHMIEL LEVINE, M D

Director of Metabolic and Endocrine Research, Michael River Assertion



THE UNIVERSITY OF CHICAGO PRESS
CHICAGO ILLINOIS

THE UNIVERSITY OF CHICAGO COMMITTEE ON PUBLICATIONS IN BIOLOGY AND MEDICINE

LESTER R DRAGSTEDT - R WENDELL HARRISON FRANKLIN C MCLEAN . C PHILLIP MILLER THOMAS PARK . WILLIAM H TALIAFERRO

> PALMA ABRAHAM SOSKIN ANNE GUSSACK LEVINE

The University of Chicago Press . Oinc. Agent Cambridge University Press . London

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PREFACE

HIS volume is intended to serve as a correlative text for the teaching of carbohydrate metabolism to students of physiology, biochemistry, and medicine. If the authors have succeeded in their endeavor, they will have satisfied a hitherto unmet need in this field. The various aspects of carbohydrate metabolism usually bave been taught as separate subjects by the different departments of universities and medical schools. This can hardly be avoided under the present system of teaching organization, but the arrangement has obvious disadvantages. Not uncommonly the net result for the student is a disjointed, incomplete, and often contradictory presentation of the subject as a whole It is the hope of the authors that the use of this text as a common meeting ground by the appropriate departments of the same institution will be of bely to both student and teacher.

A fortunate corollary of this integration of the subject is that it should make the volume useful to the practicing physician who seeks to keep abreast of the fundamentals upon which his clinical applications are based. The material is not otherwise available except in an extensive and highly technical periodical literature, with which he cannot be expected to cope directly. This applies particularly to the newer knowledge of tissue enzyme chemistry and to the pathological physiology of diabetes, a subject which has undergone a revolutionary development within the past few decades

Despite its title, this volume also deals in considerable detail with certain asaspects of protein and fat metabolism. This is mentioned to emphasize the in creasingly obvious fact that the traditional didactic separation between the metabolisms of the three cluef foodstuffs is largely artificial. Those restrictions which the present authors have placed on the scope of the subject matter depend more upon their own limitations than upon any real division of the material.

The more than twelve hundred references cited by no means represent a complete bibliography of the subject. They have been carefully selected as onginal sources of crucial experimental facts or because they review certain aspects of the subject in greater detail than is feasible in this text or because they contain useful references to the many good scientific articles which could not even be mentioned in the present volume

The senior author wishes to acknowledge the major contributions of bis asso ciates, past and present, to the development of the concepts discussed in this book

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He also wishes to express his gratitude to the Michael Reese Hospital for the ample support and academic freedom granted him to the University of Chicago for the teaching and intellectual associations which he has been privileged to enjoy and to the Committee on Publications in Biology and Medicine of the University for the stimulation without which this book might not have been under taken

Acknowledgment of indehtedness is also due to a number of authors and publishers as noted in the text for permission to use certain published materials and to Lola Kupfer Reis for her painstaking work in typing this manuscript

> S S R L

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PART I THE BIOCHEMISTRY AND ENERGETICS OF CARBOHYDRATE METABOLISM



CHAPTER 1

THE IMPORTANCE OF CARBOHYDRATES IN NUTRITION

HE importance of carbohydrates in human nutrition has varied greatly at different times and in different parts of the world Grains, fruits, and vegetables are the natural foods which are high in carbohydrate content. Meat, fish, and dairy products are relatively poor in this constituent. Before the development of the modern food processing and distributing industry (and, at the present time, in those parts of the world which have not undergone this development) the proportion of carbohydrate in the doet of any region was largely gowered by the local flora and fauna. Thus, even now the proportionate intake of carbohydrate is high in tropical countnes, where vegetation is luximous and where the chimate leads to rapid spoilage of meat products. For the obverse reasons, the inhabitants of the Far North have always lived on a diet which consists chiefly of meat and fish. Adequate nutrition is possible at both extremes of this range of dietary variation, provided that the need for calonies essential food factors, vita mins and minerals is met (1, 2, 3, 4).

Although there has been some change during the last fifty years in the food sources from which the carbohydrates are derived, the proportion of carbohydrate in the detary of the United States has remained at about 50-60 per cent of the total calonic intake. Since certain foods which are high in carbohydrate content are relatively inexpensive, the proportion of carbohydrate in the diet has been greater at lower economic levels than in the more prosperous groups of the population. However, the poorer nutritional status of the lowest income groups is not so much a reflection of their high carbohydrate intake as it is a result of the particular foods from which they derive their carbohydrates. The highly refined grains and sugars, which have heen commercially developed largely because of their resistance to spoulage are the cheapest sources of calonics generally available. But they have coincidentally been deprived of most of the protective elements with which they are naturally associated, so that a casually releved high carbohydrate diet is likely to be poor in the essential ammo acids, vitamins, and minerals (5)

THE CARROHANDATES IN FOOD

The particular carbohydrates present in the ordinary American diet, the food sources from which these carbohydrates are derived, and the quantitative importance of each carbohydrate in the total intake are indicated in Table 1

TABLE 1
TYPES AND SOURCES OF CARBOHYDRATES IN THE AMERICAN DIETARY (6)

Carbohydrates	Approximate Percentage of Total Carbohydrate Intake	Chief Flood Sources	End Products of Digestion	Resurs
olysaccharder ol Indigestable z Celluloses and hemi celluloses	e .	Stalks and leaves of vegetables, outer covering of seeds Fruits		May be partially split to glucose by bac- terial action in large bowel Chemical hydrolysis yields galactose and
b) Fartally digestible I flution 2 Galactogens 3 Mannosans 4 Raffinose		Jerusalem artuchokes, omons, garlic Stasis Legumes Sugar beets	Fractose Galactose Mannose Glucose, fractose,	ansonce Digestor incomplete, further splitting by bactera may occur in large bowel
5 Pentosans		Fruits and gums	and galactose Pentoses	
c) Digestible I Starch and dextrus	8,	Grams, vegetables (especially tu bers and legumes)	Glucose	The most important group quantitative Iy Usually accompaned by some
2 Glycogen	Negligible	Meat products and sea food	Glucose	maltose
Disacharides 1 Sucrose	25	Cane and beet sugars, molasses,	Glucose and fruc	
2 Lactose	or	Maple syrup Milk and milk products	tose Glucose and galac-	
3 Maltose	Negligible	Malt products	tose Glucose	

*Calculated from the average d ctary of the middle-moons group in the United States

TABLE 1-Continued

Carbodydister	Apprenmate Percentage of Total Carbohydrate Intake	Ch et Ford Squrces	End-Products of Digestion	Remyls
Monstaccharder a) Herrote i Glucose g Fructose	50 10	Fruits, honey com syrup Fruits, honey	Glucose	In fruits and vegetables the contents of glucose and fruitose depend on species
S Galactose			Galactose Mannose }	These monosachandes do not occur in free form in foods see under lactor and mannotans.
b) Pentoses z Kibone z Xylone 3 Arabinose	000	***	Ribore Vylose Arabinose	These monosacchandes do not occur 22 free form in foods They are derived from pentonans of fruits and from the nucleic acids of meat products and sea from
rbohydrals derivations 1. Ethyl alcohol 2. Lactic scol 3. Make scol 4. Citric scol	Variable Negligible Negligible	Fermented Inquors Mith and milk products Fruits Fruits	Absorbed as such	These substances are the products or natural or induced carbohydrate breakdown

THE DIGESTION OF CARBOHYDRATES (7)

The digestion of carbohydrates starts in the oral cavity. Here the secretion of the parotid gland, which contains an amylase called "ptyalin," is mixed with the food and hegins the conversion of starch, glycogen, and the dextrins into maltose. This digestion continues in the stomach until the hydrochloric acid which is secreted there destroys the amylase activity and substitutes acid hydrolysis for enzymatic splitting. If continued long enough, the acid hydrolysis can reduce all the digestible carbohydrates to the monosacchande stage. However, the stomach usually empties itself before this can occur, and the digestion of carbohydrate is taken up by the enzymes of the small intestine, operating in the more alkaline medium which prevails there. The enzymes in the small intestine are an amylase secreted by the pancreas, and an amylase, a maltase, an invertase, and a lactase secreted by the wall of the small bowel. All these enzymes are capable of splitting the particular sugars which they attack to the monosaccharide stage.

We have accounted for the digestion of starch, glycogen, the dextrins, and the disacchardes. Those sugars which are ingested in the form of monosaccharides do not require digestion. All the remaining carbohydrates pass through the stomach and small intestine unchanged. In the large howel they are subjected to the enzy matic influence of the profuse hacterial flora which is normal there, and they may he broken down to monosaccharides to some extent. It is possible that minor amounts of carbohydrate are made available in this manner for absorption into the blood stream (see Fig. 1).

THE ABSORPTION OF CARBOHYDRATES

The monosaccharides, ingested as such or arising from the digestion of carbo hydrates, are practically completely absorbed in the small intestine Small amounts may be absorbed from the stomach. It is also possible to show that, when solutions of monosaccharides are introduced into the large bowel for experimental or therapeutic purposes, some sugar can be absorbed from this portion of the gastro intestinal tract (8, o)

Two types of absorption occur in the small intestine (a) a specific absorption of particular monosacchardes, probably involving a phosphorylation process, and (b) a non specific absorption of all monosacchardes, by diffusion resulting from osmotic forces across the mucous membrane (10, 11) Glucose, fructose, and galactose are absorbed by both processes Consequently, the absorption of these sugars differs in two respects from that of those sugars that are absorbed hy diffusion alone they are absorbed more rapidly, and their rates of absorption are largely independent of their concentrations in the intestine (12) The explanation for the greater efficiency of specific absorption is apparently the coupling of the monosaccharde with phosphate as soon as it diffuses into the wall of the intestine This phosphorylation is a rapid process, so that the gradient of the concentration

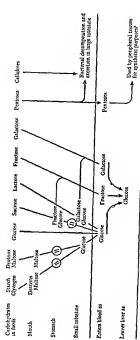


Fig. 1 - Products of carbehydiste digestion at various levels of the pastio-intential tract, and subsequent fate. © indicates that the same prod ucts as at the preceding level continue to appear

of free sugar between the lumen and the wall of the gut is much steeper than when absorption proceeds by diffusion alone

The actual rates of absorption of the three monosaccharides which are phos phorylated vary rather widely, though all are much higher than the absorption rates of such monosacchandes as mannose or the pentoses, which are handled by diffusion Thus it has been shown in rats that, if the rate for glucose is represented as 100, that for galactose would be 110, for fructose 43, and for mannose and the pentoses only o (13) There are few rehable data on the absolute rates at which the various monosaccharides can be absorbed from the gastro intestinal tract of the buman heing under normal circumstances. The best available evidence from the work of Groen (14) indicates that the rate of absorption of glucose from a 50 cm length of jejunum (small intestine) is about 8 0 gm per hour, that for galactose, about 9 5 gm per bour, and that for fructose, about 5 gm per hour These rates are for concentrations of sugar of 10 per cent and above Below 10 per cent the rate of absorption varies directly with the concentration

From the practical standpoint the figures quoted above may have little re lationship to the rate at which a monosaccharide enters the blood stream, wheth er eaten as such or arising from the processes of digestion under the usual con ditions of feeding Under the latter circumstances the time which elapses he fore it is absorbed from the gastro intestinal tract will be governed largely by (a) the rate at which it enters the small intestine and (b) the mixture of foods in the small intestine at the time of absorption. The rate at which the sugar ar rives at the small intestine depends largely on the motility of the stomach and the control of the pyloric sphincter, which can be affected by such various phe nomena as hunger, emotion, local irritation (including condiments), and the composition and consistency of the food mass after mastication (15) The food mix ture in the small intestine affects the rate of absorption by competition of the various constituents in the mixture for the absorbing surface of the mucosa and, in the case of those monosaccharides which are specifically absorbed, by competi tion for the available phosphorylating capacity (15)

Other factors which influence the amount of carbohydrate absorbed in a given individual at a particular time are (a) the normality of the mucous membrane of the small intestine and the length of time during which the carbohydrate is in contact with it. (b) endocrine function, particularly that of the anterior pituitary gland (16), the thyroid (17), and the adrenal cortex (18), and (c) the adequacy of vitamin intake, especially that of the B complex (19, 20, 21) Since the absorption of the important end products of carbohydrate digestion requires chemical ac tivity by the mucous membrane, it is ohvious that any abnormality of the muco sal cells might interfere with carbohydrate absorption Enteritis (inflammation) is a not uncommon disturbs ice of this kind Coeliac disease (22) may represent a more obscure disturbance of a similar nature However, even when the mucosa is

normal, an excessive rate of movement of the carbobydrate along the gastro intestinal tract, accompanying diarrheas of various origins, may burry a portion of the ingested carbobydrate into the large bowel before it can be absorbed

Normal absorption of carbohydrate does not occur in the presence of an antenor pituitary deficiency. This probably depends, for the most part, upon the secondary hypofunction of the thyroid gland, for the same result may be obtained after removal of the thyroid gland when the hypophysis is intact. Furthermore, the defect in absorption accompanying hypophituatism may be relieved by the administration of thyroid extract (16) Indeed, Althausen and co workers (17, 23) have attempted to make use of this phenomenon as a diruical test of the state of activity of the thyroid gland. They administer a standard amount of galactose by mouth, follow the rise of galactose concentration in the blood, and use the rate of the latter as a cirction of thyroid function in

The adrenal cortex influences carbohydrate absorption through its regulation of the sodium chloride (NaCl) exchange in the hody. The absorption of carbohydrate from the intestine is subnormal in adrenal cortical deficiency but can be restored to normal without the use of adrenal cortical extracts if the NaCl of the blood is raised to normal levels by adequate sait intake (18)

Insulin, which has such an important influence on other aspects of carbohydrate metabolism, is without apparent effect upon the absorptive capacity of the intestinal mucuus membrane

Deficiency of the B complex is associated with diminished absorption of the hexoses (19) Recent work on this subject has been concerned with the separate effects of the various pure components of the complex Thamme, panothenic and, and pyridoxine affect absorption Riboflavin is without action (20, 21)

THE DISTRIBUTION OF CARBOHYDRATE IN THE BODY ITS FUNCTIONS AND USES

In order to understand the distribution of carbohydrate in the body and appreciate its particular functions and uses, it is necessary first to consider the relation of carbohydrate metabolism to that of the other two major foodstuffs

Protein constitutes 75 per cent of the dry weight of the soft tissues of the body (24) In view of the recent knowledge as to the protein nature of the tissue en zymes, it is a fair generalization to say that the proteins, together with the bor mones, vitamins, and mineralls, constitute the metabolic machinery of the body In emergencies a certain amount of the protein machinery can be broken down and converted into fuel 'However, the amount of body protein which is available for this purpose at any one time is strictly limited as is also the length of survival

during exercise indicates that it is of secondary importance, probably to supply carbohy drate or carbohydrate intermediates. The results of experiments on fat utilization during missical work have demonstrated that this substance is used indirectly. There is no experimental evidence at the present time for the direct utilization of fat by mammalian muscle. However, the indirect utilization of protein or fat must be an efficient process since the exclusive freding of these substances to man does not have a marked effect on muscular efficiency during short periods of eversise.

The significance of the foregoing from the standpoint of nutrition is obvious If carhohydrate is not available in foods, it must be made by the body from those materials which are in the diet, in order to satisfy the fuel requirements of the active tissues. The eating of adequate amounts of carhohydrate therefore spares the body the work of making its fuel. This role of carbohydrate is naturally more

TABLE 3
DISTRIBUTION OF CARBOHYDRATE IN VARIOUS TISSUES OF RAT, DOG AND MAN
(Figures, Redresent Rapers Found on a Mixed Det)

	(-10		,	a explored and		
_	RAT		1	206	MA	и
Trasus	Glycogen (Per Cent)	Glucose (Mg per Cent)	Glycogen (Per Cent)	Glucose (Sig per Cent)	Glycogen (Per Cent)	Glucose (Mg per Cent)
Skeletal muscle Liver Heart Kidney	0 81 1 06 (33)* 2 5 -8 3 (33) 0 3 -0 6 (33)	50-70	0 55 (35) 6 10 (35) 0 47 (35) 0 15 (35)	40-6a	0 4-0 6 (35) 1 5-6 0 (37) 0 4 (37)	
Brain Skin Blood and extra cellular fluids	0 08 (34) 0 07 (39)	77 (39) 90-129 (33)	0 1 (34) 0 08 (38)	57 (34) 72 (38) 60–80	0 08 (18)	60-82 (38) 60-90

^{*} Figures la parentheses refer to hibl ngraph cat references at end of chapter

important during moderate or severe muscular evertion than when the body is at rest. The great demand for fuel accompanying muscular exercise may rapidly exhaust the cathohydrate stores. This is evidenced by a decrease in glycogen content of the liver and muscles and, if the exertion is sufficiently severe and prolonged, may result in an ahnormal lowering of the blood sugar level (41). These phenomena are accompanied by microsed breakdown of body protein (which is reflected in an increased excretion of nitrogen in the urine [40]) and by an accelerated breakdown of hody fat (as evidenced by a rise of the level of ketone bodies in blood and urine [42]). When violent exercise is preceded or accompanied by a large intake of carbohydrate, the body works somewhat more efficiently, as judged by the calories expended per unit of oxygen intake. The increased introgen excretion and ketone formation are also minimized. The latter two effects of carbohydrate examples of its protein sparing and its antiketogenic actions.

The efficiency of carbohydrote as a fuel — It has been noted above that carbohydrate is a more efficient fuel for muscular exercise than either protein or fat. This does not imply that portions of the protein or fat molecules are wasted when they

are used. It does mean that the protein and fat molecules, when used as fuel, yield less than their total calone value in the form which can be used by muscle. The remainder is used for the conversion of these molecules into suitable fuel. These conversions occur largely in the liver, which supplies the other organs with fuel by way of the blood stream.

Since the amount of glycogen present in the muscle at any one time is sufficient for only short periods of work, the carbohydrate used by the muscle must eventually come from the blood sugar. The glycogen within the muscle cells may be reasonably supposed to serve best in emergencies, when the muscle is unable to draw sugar from the blood as quickly as needed. But, as a matter of fact, glycogen is more than merely a conveniently packaged form of carbohydrate hing on the

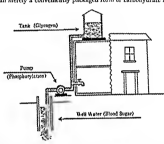


Fig. 2 - Nechanical analogy illustrating the advantage of tissue glycogen over blood sugar as an emergency fuel. (Soskin [44])

pantry shelf It is now known that more energy is derivable from a certain amount of glycogen than from an equivalent amount of blood sugar. It requires a critary amount of energy to bring the blood sugar into the metabolic system of the muscle (as herose-6 phosphate [43]), and therefore all the energy inherent in the flucose is not a salable for useful work. On the other band, the breakdown of glycogen to the same stage does not require the addition of energy and bence makes all its inherent energy quickly available (43). This is not to say that one gets something for nothing from glycogen, for some energy was required to build up the gly cogen in the first place. But this energy was expended during a quiescent period when plenty of it was available.

The above situation is analogous to that portrayed in Figure 2 (44) Here the water in the well represents the blood sugar, the pump stands for the phosphory-

lating mechanisms, and the tank on the roof represents the glycogen store. It is readily understandable that, when the tank contains stored water, the tap can deliver a rate of flow far heyond the rate capacity of the pump. The water stored during periods when the tap is closed is at a higher level than the original source of the water and also stores some of the energy applied by the pump. This poten tail energy is released when the tap is spend. Too great an outflow from the tap may, of course, exhaust the stored water and reduce the flow from the tap to the rate at which the pump is capable of operating. A similar situation may occur in muscle when excessive rates of work over prolonged periods are attempted

The application of these physiological facts to clinical phenomena is evemplified by the greater stores of glycogen and of phosphate esters found in the muscles of animals which have been trained to perform prolonged work (45). This probably also applies to the physical abilities of manual laborers and of athletes. Conversely, the characteristically low muscle glycogen levels found in poorly controlled diabetic patients and in byperthyroid individuals are accompanied by muscular weakness.

Special functions of corbohydrate in the liver —Aside from its use as fuel in the liver, carbohydrate in this organ bas protective and detoxifying actions and a regulating influence on protein and fat metabolism

The liver of a well fed normal animal contains a high percentage of glycogen, as compared to any other tissue. It is known that such a liver is more resistant to various types of noxious agents than one which has been deprived of its glycogen by starvation or disease. This has been shown in animals for such various types of poisons as carbon tetrachloride (46), alcohol (47), or arsenic (48) and in man for a variety of diseases accompanied by toxemias of bacterial origin (49, 50). The defenses of the liver against tone agents are of great importance to the body as a whole, for it is one of the chief functions of this origin to remove or destroy such towns before they reach other vital tissues which are not equipped to deal with them. From this point of view, the maintenance of a high glycogen level in the liver is an essential for the health of the whole organism.

It is now known that most of the glycogen of the liver is present in the form of a complex with protein (53) It is a reasonable assumption that, just as the protein part of the complex stabilizes the glycogen, so the glycogen would tend to protect the protein. More definite knowledge is available as regards the role of carbohydrate in specific chemical reactions which transform certain possons into relatively innocuous substances. One such mechanism is the conjugation of gly curonic acid detived from carbohydrate with possons which possess a hydroxyl group (52, 53). Indeed, this mechanism is one of the means by which the body regulates its steroid hormone metabolism and protects itself from the harm which could result from an excess of the sex hormones (54). It is also possible that the carcinogenic substances of the steroid type might be disposed of in the same man

THE IMPORTANCE OF CARBOHYDRATES IN NUTRITION ner Another hepatic mechanism is the acetylation of such substances as p aminoner Anouser nepaus mecoanism is one accommon or such substances as p anno-benzoic and (55) and sulphanilamide (56) In this type of conjugation the acetyl perizone acid (55) and surprimariamine (50) an time type of conjugation the acetyr groups are derived from carbohydrate probably via pyrinyate and acetyr phosgroups are derived from caroonydrate pronactly via pyrtivate and accetyl phosphate. The rates of glycuronate formation and of acety latton have been shown to Ι¢ phase the rates of glycuronate formation and of acets lation have ocen shown depend directly upon the concentration of carboby drate in the liver (56, 57)

epend directly upon the concentration of caronogurate in the liver 150, 57/ The protein sparing action of carbohydrate has already been mentioned. This The protein sparing action of carbonydrate has already oven mentioned. A mas action occurs partly in the liver, for it is this organ which is primarily responsible. action occurs parity in the liver, for it is this organ which is primarily responsible for the deamination of amino acids. Up to the point of deamination the fate of for the deamination of amino acids. Up to the point of deamination the fate of amino acids in metabolism has not been finally determined. They may be used as annus acuta us metauousm nas not ucen mrany determined. A ney may ne used as building blocks from which to form proteins for the repair or growth of tissues, or building blocks from which to form proteins for the repair or growth or tissues, or they may be broken down for use as fuel. Once deamination has occurred, the oney may be oroxen gown for use as fuer once destination may occurred, the amino acids are divorced from protein metabolism. The amino group is converted amino acids are divorced from protein metabousm. The amino group is converted to use and excreted, while the non nitrogenous fraction is either used as a source to use and exercised while the non nitrogenous fraction is either used as a source of energy or converted to carbohydrate or fat 'The rate of deamination in the liver or energy or converted to carbobydrate or latt the rate or deammation in the liver decreases as the available carbobydrate increases. An ample supply of carbobyoccreases as an available carbooyutate increases. An ample supply of carboby-draft thus conserves the products of protein breakdown in a form which may be arate thus conserves the products of protein oreascown in a form which may be used by the body to build or maintain its own protein structure. To put it in an used by the body to build or maintain its own protein structure. Lo put it in an other way, a minimal intake of protein which may be adequate for the body's oner way, a minimal intake of profein which may be adequate for the body s needs when taken together with good amounts of carbohydrate, may become in adequate when the carbohydrate intake is deficient (58)

ucquare when the carponyurate intake is denoted; (50)

The availability of carbohydrate to the liver also determines bow much fat is and availability of carbonyurate to the aver also determines only much tar is broken down by this organ. There is no direct index of the rate of lat metabolism. oroxen cown oy uns organ there is no circu mack of the rate of lat metabodism in the liver, for, unlike protein metabodism fat metabodism is not accompanied by in the river, for, untike protein mecanousm far metanonism is not accompanied by the excretion of a characteristic end product in the unne. However, it bappens the exercison of a characteristic end product in the unine storecter, it completely metabolized by the liver and that the end that tatty acids are not completely metabolized by the liver and that the end products of fatty acid metabolism in this organ are the so-called ketone bodies? Products of latty and metabousm in this organ are the so-caused ketone bodies."

A hydroxybutyric and acetoacetic acids (59 60, 61) These ketone bodies must p nydroxyoutyric and acctoacetic usus 159 to, o.). These actoric volume invastities go to the peripheral tissues for complete oudation. Ordinarily the rate of then go to the peripheral dissues for complete obtained Urumanily the rate of breakdown of fat and of the formation of ketone bodies is such that the latter are oreascown of lat and of the formation of actions bodies is such that the latter are promptly disposed of by the peripheral tissues, so that no significant amounts ap Prunpity suspessed of by the peripheral tissues, so that no significant amounts ap pear in the blood or unne. But when fatty acid breakdown becomes excessively read to the proof of clinic but when rate) acid breaknown becomes excessively and the rate of ketone formation in the liver begins to exceed the rate of dis iapso and the rate or section formation in the over pegins to exceed the rate of dis posal by the peripheral fissues, there begins to occur an accumulation of the ke posal by the peripheral tissues, there begins to occur an accumination of the ke tone bodies in the blood and an excretion of these substances in the time (ketosis) tone nouse in the nigog and an excretion of case souscauces of the utility (kerosis). Under these circumstances in an otherwise normal animal the administration of Vauer tinese circumstances in an ometwise normal animal the administration of carbohydrate causes a prompt disappearance of the ketone bodies (antiketogenic catronyurate causes a prompt disappearance of the actions owness (annactogenic action). This effect of carbohy drate occurs in the liver and is due to an inhibition oction) this enect or caroon) urace octurs at one user and so one to an industrion of the breakdown of fatty acids. Together with the protein sparing action of or the oreastown of taily acids. Agreemer with the protein sparing action of carbohydrate, its antiketogenic action serves to regulate the proportion of the an amino acid (chap u p 39)

Under certain circumstances the non nitrogenous fraction may also be reminated and restored as

different foodstuffs which are prepared by the liver for use as fuel by the peripher al tissues

In discussing the special functions of carbohydrate in the liver we have referred both to its "glycogen content" and to the "availability" of carbohydrate to this organ. These terms may or may not be synoupmous, for it is still not known whether sugar may be used directly by the liver calls or must first be built up to glycogen. In any case, the glycogen content of the liver is a good index of the amount of carbohydrate which is available to the hepatic cells, and from a nutritional standpoint it is important to remember that carbohydrate is the foodstuff which leads to the highest levels of liver glycogen. Fairly good glycogen stores in the liver can be obtained when protein is predominant in the diet, while a high fat diet results in a liver which is poor in glycogen (62, 63). The medical uses of the high carbohydrate duet or of the intravenous administration of dextores solution are directed toward the protection of the liver by insuring rich glycogen stores (50). Protein has been used with the same ultimate purpose in mind, but it is less effective, probably in proportion to its convertibility to surgar.

Carbohydrate and the heart - The previous discussion of carbohydrate as the most efficient fuel of muscular exercise, and of the muscle glycogen as an important emergency source of contractile energy, applies in even greater measure to cardiac muscle than it does to skeletal muscle. The latter can in some measure accommodate itself to a decreased supply of carbohydrate by decreasing its work. The heart cannot stop to rest A temporary reduction in the supply of sugar to the normal heart (as in induced attacks of bypoglycemia) has little apparent effect on the organ, although a definite change in the electrocardiogram may be noted (64) The apparent lack of influence of bypoglycemia on the normal beart may be due to the good glycogen stores to be found there But, in the heart which is damaged by disease and in which the initial glycogen stores are poor, hypoglycemia may precipitate stenocardial symptoms with angina and may even result in death This has been noted for diabetic (65), as well as for non diabetic, cardiac patients, and in both it has also been observed that they may do better when the blood sugar is somewhat elevated even above the normal range. High carbohydrate therapy has been successfully used on this basis (66)

The indispensability of carbohydrate to the central nervous system — Of all the organs and tissues in the body, the central nervous system is most dependent upon the minute by minute supply of glucose from the blood. In connection with the discussion on the fuel of muscular exercise it was stated that carbohydrate was of primary importance, while protein and fat could be used only indirectly. As regards the central nervous system, it has been well established that only carbobydrate can be used (67, 68, 69). The need of nerve tissue for glucose is even more specific than the previous statement would indicate. It is true, when slices of brain

tissue are studied in vitro regarding their ability to maintain respiration at the expense of various substrates, that a number of degradation products of glucose will serve as well or better than glucose itself (67) However, none of these inter mediates have been shown to have any ameliorating effect upon the hypoglycemic symptoms caused by lowering the blood sugar level in vivo (70) In other words, plucose as such bas a specific influence and is indispensable for the maintenance of the functional integrity of the nerve tissue. When the blood sugar is lowered in a living organism, those tissues which have ample stores of glycogen may use the latter to tide them over the lean period. The nervous tissue has little glycogen, and it is doubtful whether the little which is present can be mobilized for use in emergencies. The glycogen content of nervous tissue remains more or less con stant under most conditions, including hyperglycemia and hypoglycemia, and may be largely an integral part of the nerve structure (34) The unavailability for metabolic use of the glycogen present in the nerve cells is evidenced by the dramatically rapid development of hypoglycemic symptoms when the blood sugar is lowered

THE TRANSFORMATION OF CARROTYDRATE INTO FAT

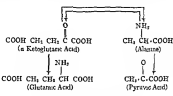
In the previous discussion of fat as a fuel storage material it was pointed out that, when food in excess of calonic expenditure is ingested (whether in the form of carbohydrate, protein, or fat), the equivalent of the excess calonies is deposited as fat in the adipose tissues. With this in mind, it is, strictly speaking, incorrect to label any of the foodstuffs as being particularly "fattening". Any one of them can be so if taken in sufficient quantities. But because of its proportion in the diet, its lower cost, and its use in confections, carbohydrate is quantitatively the most important precursor of fat.

The fat which anses from carbohydrate in the body is the so-called "hard" fat, composed, in the main, of the highly saturated plantitie and stearne acids (71). This is probably of more concern to stock raisers than to human nutritionists. The former have long known that they could control the physical qualities of the fat in meta by varying the proportion of carbohydrate and of oils in the diet of their animals. Of course, carbohydrate cannot completely substitute for fat in the diet, since it does not carry with it the essential fatty acids and the fat soluble vitamins, which cannot be manufactured by the body.

THE INTERPELATION OF CAREOUNDRATE AND PROTEIN METABOLISM

Earlier writers on metabolism have talked somewhat loosely of the formation of protein from carbohydrate Strictly speaking, such a transformation does not occur, because the amino groups which characterize the building stones of proteins are derived from amino acids or proteins which are ingested as such Schoen-

heimer (25) has demonstrated that, when ammonium salts are ingested, the NH, may combine with carbohydrate derivatives to form amno acids. But what ordinarily occurs is the exchange of the amino group of an amino acid with the keto group of a keto acid (derived from carbohydrate), a process known as "transam nation" (72, 73). In this process the carbon residue of the amino acid reverts to a carbohydrate intermediate, so that there is not necessarily any quantitative in crease in the amount of protein procursor resulting from the reaction. What the body gains from the interchange is the ability to transform one amino acid, which it may have in excess to another, which it may need. For example, by exchanging with a ketoglutarate, alanine may be transformed to glutamic acid, with pyruvic acid as the by product (Fig. 3).



Fic 3 -Example of transamination

THE IMPORTANCE OF THE VITAMIN B COMPLEX IN CARBOHYDRATE NUTRITION

It is now known that many members of the vitamin B complex play an integral part in carbohydrate metabolism and that the requirement for this group of vita mins depends upon the amount of carbohydrate which is eaten. Since this is so, why did not the knowledge of its existence arise much earlier in human experience and why did not the race suffer from the lack of such knowledge? The answer to these questions lies in the fact that it is only in comparatively recent times that the natural union between the vitamin B complex and carbohydrate, which exists in whole grain and plants has been broken by the industrial processing of foods Before this occurred, the supply of the B vitamins was automatically adjusted to the amount of carbohydrate eaten, so that the occurrence of vitamin B deficiency, with its consequent disturbance in nutrition, is a comparatively recent development in the Western world. In the Orient the earlier large scale introduction of polished rice led to the first known instances of vitamin B deficiency (deribert) and, indeed to the first known instances of vitamin B deficiency (verbert).

The vitamins, as the name signifies, were first regarded as mysterious elements,

essential for life. As the different vitamins were successfully recognized and extracted in concentrated form from their natural sources, experimentation with these products led to the recognition of definite clinical syndromes resulting from their lack and cured by their administration. More recently the actual chemical identity of many of the vitamins has been established, and a number of them have been synthesized. Coincidentally with the latter events, the development of issue-enzyme chemistry has revealed a great deal about the chemical steps through which the foodstuffs are broken down and used for energy. It is now known that each of the chemical steps is accomplished by the activity of one or more enzymes (protein catalysts) and that each of the enzymes requires one or more oriactors for its optimal activity. In some instances the cofactor is a simple numeral substance, like iron or magnesium or phosphorus, in other cases the cofactor is a more complex organic substance, known as a "coenzyme". Thus far, those vitamins whose functions are known bave been found to be coenzymes or to give nie to coenzymes in the body (75).

Figure 4 outlines the known steps in the breakdown of carbohydrate and indicates the points at which the various components of the vitamin B complex play an essential role. The role of various minerals in carbohydrate metabolism is similarly indicated. It may be seen that definite knowledge is available regarding only three B factors, namely thiamine mectunic acid, and riboflavin. It is to be expected that similar functions will eventually be found for the other factors in the B complex.

Since the breakdown of carbohydrate is essentially similar in all tissues and organs, it follows that a vitamin B deficiency will impair carbohydrate metabolism in every structure of the body. The clinical syndromes which have been described are, therefore merely the most obvious manifestations occurring in those itsues and organs to at suffer most acutely and that are most easily accessible to examination. Consideration of Figure 4 also shows the fallacy of regarding any single factor of the B complex as more important than another, for the normal chain of events can be broken by a lack of any one of them. For this reason and until we have itsolated and know the precise function and optimal proportion of each component part of the B complex, a natural source containing all the factors remains the best protective dietary supplement with which to avoid the evils of modern food refinement.

THE UTILIZATION OF SIMPLE SUGARS OTHER THAN CLUCOSE

In the previous section on the distribution of carbohydrate in the body it was pointed out that all the hexoses absorbed from the gastro-intestinal tract are converted into either glucose or glycogen. This conversion, which takes place largely in the liver, is ordinarily so efficient that there is little need to consider any other fate which signars like fructose and galactose may undergo However, under special

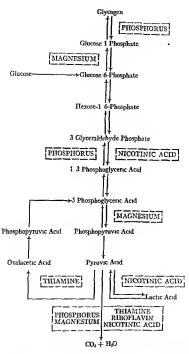


Fig. 4—Points of action of vitamins and minerals in carbohydrate metabol sm. The substances reurred for a particular reaction are necessary in both directions of the reaction

circumstances when the function of the liver is impaired or when these sugars en ter the blood in overwhelming quantities there occur interesting anomalies of carbohydrate nutrition which deserve some brief mention. Lactose is also of interest because of its formation in large quantities by the lactating breast of the female at which time it may appear in the blood and the urine. The pentoses are sometimes movived in a hereditary anomaly of metabolism

a) Fructors —While the conversion of fructose to glucose occurs largely in the liver, there is some evidence that it may take place to a smaller extent in the in testinal mucosa and the kidney (28-29-76). Recent work indicates that there are probably two chemical pathways from fructose to glucose in the liver. The fructose may be phosphate vide to fructose 6 phosphate which is converted to glucose 6 phosphate and then split by the liver phosphatise to yield glucose (77). The phosphorylated fructose also appears to be more readily degraded to lactic acid than is glucose 6-phosphate. Hence when fructose appears in excess in the blood it is accompanied by a rise in lactic acid (78). Some of the latter may be converted to glucose or elyocogen by the liver.

When any of the foregoing hepatic mechanisms are impaired either by liver as a property of the foregoing hepatic mechanisms are impaired either by liver culty in disposing of the fructose taken in through the gastro-intestinal tract and it accumulates as such in the blood (79). Since it is a substance which is not beld back by the kidney as efficiently as is glucose it appears in the urine in abnormal quantities. Fructose is a reducing sugar which is not distinguished from glucose by the routine chemical tests. From the medical standpoint it is therefore important not to confuse fructouria with diabetes mellitus.

b) Lactose and galactose —Lactose is split into glucose and galactose in the process of digestion. It may therefore he considered together with the galactose which is ingested as such. However, the presence of lactose in milk and milk products renders it much more important than galactose from the nutritional standpoint Lactose also has the special virtue of altering the intestinal flora in such a manner as to produce a more acid environment, which favors the more complete absorption of ingested calcium (86).

There is some recent evidence that suggests that galactose is converted to glucose in the liver hy phosphorylating steps similar to those described for fructise (81). Little beyond this is known For example the lactating breast manufactures lactose and presumably has galactose available for the purpose (82) but it is not known whether all the galactose is made in the breast or whether some of it ofiginates in the liver and is transported to the breast. Both lactose and galactose may be found in the blood and urine of lactating females so that the mere presence of these abnormal constituents does not give any indication as to their site of origin As with fructose it is of importance medically to distinguish between galactosuria and factosuria and glucosuria.

22 In the previous discussion of the special functions of carbohydrate in the liver

mention was made of its protective and antiketogenic action. Liver glycogen that is formed as a result of the intake of galactose or of lactose may perbaps be more beneficial to the organism than glycogen that originates from other materials This is because, for some unknown reason, the "galactose glycogen" is more stable It has been shown that, when galactose is administered to animals together with a ketogenic agent, the ketosis which follows is less than when glucose or fructose are similarly administered (15)

c) Pentoses -In contrast to the hexoses, which are important energy materials the five carbon atom sugars are much more important as part of the machinery of the body Pentoses are incorporated in at least one vitamin (riboflavin), several tissue coenzymes (diphosphopyridine nucleotide, triphosphopyridine nucleotide, and alloxazine adenine dinucleotide), and all the nucleoproteins. However, when pentoses as such are ingested, they are not utilized but are eliminated, more or less quantitatively, in the urine and feces. It is possible that the pentoses which are eaten in combined form as part of natural food constituents (riboflavin and the nucleotides, for example) do contribute to the pentose content of the tissues It is known that the body is able to synthesize pentoses for itself from glucose by way of glycuronic acid (83) The bereditary anomaly known as "essential pento suria" is as yet unexplained

SUMMARY

We have seen that carbohydrate is not only the primary fuel of the body but is also involved in important portions of its functional machinery. The carbo hydrate stores, though relatively small as compared to fat, play a protective role in some of the most vital organs. They may be of the utmost importance when a rapid source of energy is required, to enable the organism as a whole to cope with an emergency in its environment Despite all this, however, the evolutionary processes have resulted in so flexible a metabolic system that the higher mammals and man can get along very nicely when little or no carbohydrate is available Under these circumstances the body makes its own carbohydrate fuel from non car bohydrate materials. But this is a wasteful process, because some energy must be used for the conversions, and there is more wear and tear of the metabolic ma chinery

If, with the foregoing considerations in mind, we could divorce ourselves from previous dietary experience and were to attempt to construct an ideal adult diet, we would choose the following

Protein sufficient in quantity and quality to repair the protein machinery from day to day, and a little extra, to be on the safe side. In the same category we would place a sufficiency of all the vitamins and minerals

2 Enough fat to carry the essential fatty acids and fat soluble vitamins and to make it unnecessary to eat too large a bulk of other food

3 Carbohydrate sufficient to supply all the rest of the calones necessary to

maintain weight

The det which has been outlined is a fair approximation of that which the human race has actually adopted on the basis of experience, in those fortunate parts of the world where food resources are not and the choice is not limited (84)

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CHAPTER 11

THE ENZYMATIC MACHINERY OF CARBO-HYDRATE METABOLISM

N THE process of digestion or in the liver after absorption, carbohydrates are largely converted to glucose Hepatic gluconeogenesis leads to the same end product. The further course of carbohydrate metabolism is therefore chiefly concerned with the chemical transformations undergone by glucose. These in clude the synthesis of glycogen and the formation of fat. But more basic than either of these is the hreakdown of the sugar to carbon dioxide (CO₂) and water (H₂O), with the liberation of the energy that supports the various functions of living cells

Lavoisier's analogy of the burning candle introduced the concept of oxidation in the living organism and the use of the term "combustion" to describe the ultimate breakdown of foodstuffs in the body. The analogy was apt and useful at the time. The living organism, like the hurning candle, required oxygen and produced CO, and H₂O What could be more natural than the conclusion that the lungs served as a furnace, where the inspired oxygen united with carbon and hydrogen from the blood to produce beat, energy and the appropriate end products (1)? During the first half of the nineteenth century the discovery that the blood con tained O2 and CO2 resulted in a shift in the location of the theoretical furnace from the lungs to the blood (2) However, the development of histological and biochemical techniques soon led to the realization that the individual tissue cells were the functional units of metabolism, while the blood served mainly as a medium of transport (3) This, in turn gave birth to the vague and somewhat vitalistic conception of the ability of the body tissues to 'oxidize" food materials and to derive heat and energy therefrom At that time, the word "oxidation" was not used in the strict chemical sense of today. As then used, it meant the simple addi tion of oxygen to molecules or carbon fragments of the original foodstuffs within the tissue cells, and the liberation of energy by complete oxidation of the food stuffs to CO2 and H.O This concention, with little modification, has been carried forward in some writings to the present day

The work of Pasteur on yeast fermentation initiated a series of scientific de velopments, which at first were apparently unrelated to the above but which eventually merged completely. The epoch making discovery by Buchner (4) that a cell free extract of yeast could substitute for the living cell in the process of fer mentation showed that what had been considered to be a process inseparable from

tie is, after all, only a special kind of chemical reaction—a reaction that is catayized by a complex organic substance (enzyme) in the cell. This paved the way for
n rational and materialistic explication of cell processes. Other enzymes were dis
overed and isolated. Evidence mounted that the chemical machinery of the liv
ng cell consists of a series of organic catalysts which operate on complex molecules, step by step, to produce simpler and more labile products. It was realized
that the enzymes made possible such chemical reactions in the cell as would otherwise require high temperatures or strong reagents incompatible with life. The
step by step catabolism controlled by the multiple enzymes also offered a reasonable basis for the regulated release of energy in small units, a process which
was much more reasonable, from the point of view of the use of such energy, than
the explosive type of reaction, implied in the idea of "combustion"

By the early years of this century biochemists and physiologists using hiochemical methods had collected a great deal of data concerning the kinds and amounts of intermediate metabolites present in the different tissues of the body under a variety of conditions. These data guided the enzyme chemists in the isola tion and study of the enzyme systems which were responsible for the various prod ucts. The last ten to fifteen years have winessed a tremendous and constantly accelerating growth in the application of enzyme chemistry to metabolic problems. It has become evident that, in the process called "oudation" in the tissues, molecular oxygen does not interact directly with the foodstuffs (5, 6) and that CO, largely anses by a splitting-off of carboxyl groups from lower metabolic intermediates (7). It is with these and other fundamental enzyme reactions that the present chanter will deal.

NATURE OF CELL ENZYMES

The enzymes in the living cell resemble the known inorganic catalysts in that they are more or less specific for a particular chemical reaction or type of reaction, also, in that they are not measurably consumed by the reaction which they ac celerate All the tissue enzymes which have thus far been isolated and sufficiently punified that their essential natures are known bave turned out to be proteins (8, 9). As more and more of the enzymes have been recognized and studied, it has become less possible to distinguish between purely structural proteins constituting, as it were, the skeleton of the cell (10), and the enzyme proteins, representing the active organs of the cell in fact, a tabulation of the number of enzymes present in skeletal muscle and a calculation of the proportion of the total cell protein which enzymes must represent leaves little or no room for the presence of any purely structural proteins (Table 4) (6 m.).

Studies of the optimal conditions for the activity of various enzyme proteins have uncovered a number of other normal constituents of the living cell which must be present if a particular enzyme is to exert its fullest effect. In some in

stances these accessory substances are simple ions, like phosphate or magnesium, and are referred to as "cofactors" of the enzyme. When the accessory element is a complex organic but non protein substance, it is known as a "coenzyme" (12) A protein enzyme (or the activating protein) together with its particular coenzyme and/or other cofactors is known as an "enzyme system".

THE ENZYME SYSTEMS INVOLVED IN CARBOHYDRATE METABOLISM

The following is a list of the various types of enzymatic reactions which are known to be involved in the breakdown and synthesis of carbohydrates in mam malian tissue. The enumeration is followed by a brief description of the nature of

TABLE 4

PROPORTION OF THE MUSCLE PROTEIN ACCOUNTED FOR BY A FEW
OF THE MANY KNOWN EXTRES SYSTEMS*

Catalytic System	Percentage of Total Prottin	Reference
Adenosinetriphosphatase (myosin)	50-60	Figethardt (11)
Zymohexase (myogen)	2	Herbert (102)
Lactic debydrogenase	0 4	Straub (15)
Cytochrome C	0 09-0 3	Stotz (92)
Myoglobin	0 5 -2 0	Milhkan (103)

*There are at present forty add topal known earyme systems in the muscle cell [6]. Thus relative concentrations are unknown. It is avident however, that practically all of the cell proteins are constructed of access catalytic systems.

each reaction and an important example of each type, including mention of the coenzymes and cofactors involved

- 1 Oxidation (oxidoreduction)
 - 2 Decarboxylation (oxidative and non oxidative)
- 3 Carbon dioxide assimilation (addi-
- tion of CO₂)

 4 Phosphorylation and phosphorolysis
- s Intramolecular phosphate transfer
- 6 Deamination
 - 7 Amination 8 Transamination
 - g Hydrolysis

r Oxidation —The term "oxidation" may be applied to a reaction when there is (a) the addition of oxygen atoms to a substance, (b) the removal of hydrogen atoms from a substance, or (c) the removal of electrons from a substance (13, 14). The transformation of lactic to pyruvic acid is such a reaction and may be indicated as follows.

r mol lactic acid - 2 hydrogen atoms = 1 mol pyruvic acid

The hydrogen is not given off in gaseous form but rather in the form of hydrogen ions and electrons. This means that for each hydrogen ion one electron is also released. The correct chemical notation for this reaction is therefore.

Since this particular obudation consists of the removal of hydrogen atoms, it is often referred to as a "dehydrogenation"

Lactic acid, dissolved in H,O and with free access to oxygen at 37°5 C, will be oxidized to pyruvic acid at such a slow rate as to be hardly measurable. But when a specific protein derived from animal or plant cells is added to the solution, significant amounts of pyruvic acid appear in a matter of minutes (15). This influence of the activating protein or enzyme may be regarded as one which loosens the bonds joining the two hydrogen atoms to the second, or aC, atom of the lactic acid molecule. More accurately stated, the activating protein changes the form of the electron energy, uniting the hydrogen and carbon in such a way as to increase the tendency of the hydrogen atoms to fly off (10). Thus any suitable chemical substance which can hind the hydrogen atoms (hydrogen acceptor) will remove the "loosened" hydrogen from the orbit of the lactic acid, leaving pyruvic acid (1 ig. 5) (17, 18, 10).

The hydrogen acceptor necessary for the above reaction is diphosphopyndime nuclouide (DPN) (Tig. 6) (i.) This, then, is the coenzyme which, together with the protein, makes up the lactic acid oxidase (or dehydrogenase) system Despite this nomenclature, however, the system is reversible and will actually reduce pyravia caid to lactic acid under the proper conditions (i.) The direction of the reaction depends largely on whether the DPN is present in its oxidured or reduced form (as DPN or as H.DPN) which, in turn depends upon whether other ystems when an remove the hydrogen from DPN are present (to, i.) For example, the activity of the lactic acid oxidase system in the living animal is most frequently of served during relative or absolute amoust in skeletal muscle when the H.DDN cannot readily be reoxidized and hence serves to convert pyruvic acid to lactic acid. In chemical notation the reaction may therefore be represented somewhat more completely, as follows:

While the activating protein of the lactic acid oxidace system is completely specific for the one substrate, lactic acid, and is just as specific for the particular transformation of lactic acid which we have described, the coenzyme is less discriminating. It also serves as a hydrogen acceptor for other reactions (see Table 5) Each of these reactions is catalyzed by a separate activating protein in combination with DPN Some hological oxidations are carried on by systems consisting of proteins and TPN (see legend to Fig. 6). These two groups constitute the class of pyridinoprotein enzytices (27, 23). Another group of candation systems are known as the "yellow enzymes—proteins combined with alloxazine derivatives (Fig. 7), which are yellow in aqueous solution (24, 25).

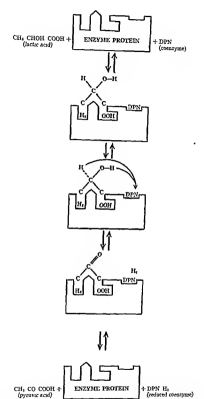


Fig. 5—A schematic representation of the configuration of an engine prote in (maginary) showing the mainer in which it is thought to anchor the substrate and the commyne and to facil tate the interaction between the free groups of both

The various oxidation systems that have been listed are responsible for the removal of hydrogen from all substrates and intermediate substances whose metabolic fate is known. The hydrogen removed from the original owner, while under the influence of a specific protein, is simply transferred to the coenzyme of the system, be it DPN, TPN, or an alloxazine. It will be noted that no mention has been made of the appearance of ovyren upon the scene As a matter of fact, the

FIG 6—Diphosphopyndine nucleotide (DPN) H = hydrogra atoms from substrate (Imphosphopyndine nucleotide (IFN) differs from DPN in possessing an additional phosphate group between the rhose units of the molecule;

TABLE 5

OXIDOREDUCTION REACTIONS AND THE COENTIMES OF ERATIVE IN THEM

Reset on	Coenzygne	Reference
Lactates: Pyruvate Acolotics: Acidebyte Bydium; phutprates: Acotoacetate Glacone: Glaconic and Malates: Oxidertate	DPN DPN DPN DPN DPN	Straub (2g) Lutwak Mann (26) Green (27) Harrison (28) Das (29) Green (20)
		* ·
		•
Vanthuezi Urc acid Aldehydesat Acids Fomarate→Succupate	Flavio Flavio Flavio Flavio	Hazs (38) Ball (39) Booth (40) Gordon (41) Fischer (42 43)

hydrogen seized by the coenzyme is passed on through a series of other systems in the manner of a bucket brigade before it finally arrives at the point where it may combine with oxygen to form H₂O. This will be discussed in detail later (p. 41)

2 Decarboxylation — Carbon dioxide is one of the end products of the complete breakdown of foodstuffs. It is not formed as was formerly thought by the direct oxidation of the carbon by molecular oxygen but arises from the splitting off of carboxyl groups (-COOH) from intermediate organic acids which arise in the course of catabolism (7). The exact mechanism of decarboxylations is a syet ob scure but we can distinguish two types the oxidative and the non oxidative. In the first of these the CO, is split off a molecule while at the same time hydrogen.

Fig. 7 - Alloxaz ne adenine dinucleotide (flav n) H = hydrogen atoms from substrate

atoms are removed from another group in the same substance. For example pyruvic acid CH, CO COOH containing three carbon atoms is oxidized to acetic acid CH₃ COOH which contains only two carbon atoms the third having been split off as CO, (44 45). In chemical notation this double process of oxida tion plus decarboxylation can be presented as follows.

$$\begin{array}{c} O \\ CH_{\begin{subarray}{c} CCOOH + H_sO \rightarrow CH, \\ CH_{\begin{subarray}{c} COOH - (2H^+ + 2) \rightarrow CH, \\ OH \end{subarray}} \begin{array}{c} OH \\ CCOOH - (2H^+ + 2) \rightarrow CH, \\ OH \end{subarray} \begin{array}{c} C+CO, \\ C+CO, \\ C+CO, \\ OH \end{subarray} \end{array}$$

In the second type of decarboxylation there is no concurrent oxidation Again

using pyruvic acid as an example this type of decarboxylation proceeds as follows (46, 47)

Just as in the oxidations the various decarboxylations are catalyzed by specific activating proteins and the process is aided by coenzymes and cofactors. The co enzyme needed for the decarboxylation of pyrivic acid is diphosphothiamine (also called "cocarboxylase") (Fig. 8) Magnesium ion is also an essential component as a cofactor in the foregoing systems (46, 48, 49).

Fig. 8 -D phosphothismine (cocarborylase)

Although the splitting off of CO, is not so well understood a process as is oxidation a number of substances are known to undergo this process (Table 6) It seems quite definite that in all cases the production of metabolic CO, proceeds in the fashion detailed for nyrunce and

TARLE 6

DECARBOXYLATIONS IN INTERMEDIARY CARBOHYDRATE METABOLISM

Pyruvate → Acetate + CO,
Pyruvate → Acetate + CO,
Pyruvate → Acetylmethylactionol + CO,
Pyruvate → Acetylmethylactionol + CO,
Pyruvate + CO,
Pocalacetate → Pyruvate + CO,
Pocalacetate → Ecologitarate + CO,
Pocalacetate → Succenate + CO,
Pocalacetate → CO,
Poc

3 Carbon durade assimilation—It has been known for some time that CO, produced by the dissimilation of foodstulls may combane with hemoglobin (ear hamino compound) (53) or may be used for the production of urea (54). It was supposed that by these and other means all the CO, produced by the mammalian organism was eventually excreted by the lungs and the kidneys. Only plants or certain autotrophic bacteria were thought to possess the ability to incorporate CO, into usable cell products (51). In 1936 this ability was first observed in bacteria (55 56), later it was confirmed for mammaliant usue (especially liver) (57, 58) that certain in vitor reactions undergone by compounds containing three car bon atoms (the troses) could be speeded up if CO, were present in the medium. It was shown that this was not a consequence of the mere presence of CO, but that the CO, took part in the reactions and was incorporated into other substances (51, 58).

,

Again pyruvic acid will serve as a good example. In the presence of the specific proteins, diphosphothamine, inorganic phosphate, and magnesium ion, pyruvic acid (a three carbon atom compound) and CO, will form oxalacetic acid (a four carbon atom compound).

This is probably the first step in the series by which pyruvic acid (or lactic acid) is reconverted to sugar and glycogen (50, 60, 61)

The use of CO, for synthetic purposes by the mammalian cell is only now being studied in detail. But it has already taken on tremendous significance, since it completely reverses the hitheric firmly accepted view that CO, is merely a waste product of animal metabolism (7, 51). It particularly affects our outlook on in direct calonmetry (p 96).

4a Photphorylation —Early in the development of our knowledge of the en zymatic breakdown of carbohydrates it was shown that the presence of phosphate was necessary for the fermentation of glucose by yeast extracts (62) and for the breakdown of sugar that takes place in active muscle extracts (63). It was later demonstrated that the phosphate is used for the formation of various intermedianes of carbohydrate breakdown which were shown to contain phosphate in their molecules (63–64). Among such metabolites are the glucose and fructose mono phosphates fructose diphosphate, glyceraldehyde phosphate, etc (cf. p. 50). The role of these phosphorylated intermediate substances in facilitating certain reactions and in the transfer of energy from one chemical reaction to another bas only recently been clucidated. We shall discuss these aspects in detail in the section dealing with the utilization of metabolic energy (chap iv, p. 60). For the present it will suffice to present the mechanics of phosphorylation by suitable examples.

The first step in the series of reactions by which sugar enters the metabolic cycle of the cell is the addition of phosphate (P) to the sixth carbon atom of the glucose molecule (05 66). The enzyme necessary for this initial reaction in animal insistes has not yet been purified, but it apparently activates the glucose molecule in such a way that it can receive a phosphate from a suitable source. The phosphate donor in this case is adenosine triphosphate (ATP) (Fig. 9), which is the coenzyme of this phosphorylation reaction. In chemical notation the reaction may be represented as follows.

The coenzyme ATP has two phosphate groups, which can be split off easily in the presence of the suitable enzymes (67, 68)

But the amount of ATP present in the cell at any one time is very small as compared to the amount of material to be phosphorylated. Hence ADP and AA must be continuously reconverted to ATP (p. 6) in order that the latter can serve as a continuous phosphate donor. The central position of this adenylic system for receiving and donating phosphate groups is illustrated in Figure 10, in which the direction of the arrows represents the direction of phosphate transfer.

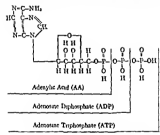


Fig. 9 -The coenzyme system for phosphory lations

ab Phaspharolists—Gly cogen is a complex molecule consisting of glucose units connected to one another by glucoside (C-O-C) inkages. Two types of inkages occur, the 1 4 and the 1 6 (69 70), as illustrated in Figure 17. The gly cogen complex is therefore, not a strught-chain polymer but a bighly branched structure. The breakdown of glycogen to heaves units is accomplished by two en zymes each of which is specific for one of the inkages. The better studied and now purified system is the 1 4 enzyme, known as "gly cogen phosphorylas." (71, 72) in the presence of inorganic phosphore and (H,PO), cleaves the glucoside inkage, leaving H,PO, attached to carbon atom x of one glucose unit and H attached to carbon atom 4 of the next glucose unit. This is analogous to a hydrolytic cleavage (H OH) except that, instead of elements of H,O, those of the orthophosphate are

added Because of this analogy the name "phosphorolysis" (compare with hy drolysis) is given to this type of reaction (104, 105, 106). The reaction is visualized in Figure 12. The 1.6 linkage is probably broken in a similar manner by the 1.6 phosphorylase (70, 72).

Phosphorolysis is reversible. The direction of the reaction is determined by the relative concentrations of glucose x phosphate and inorganic phosphate, so that removal of inorganic phosphate favors glycogen synthesis, while addition of in organic phosphate hastens glycogen breakdown (7,5,74). There is evidence that this is one of the regulating devices of glycogenolysis in the living cell.

PHOSPHATE DONORS

PHOSPHATE ACCEPTORS

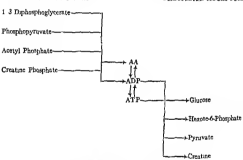
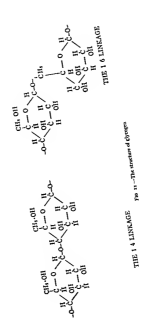


Fig 10 -Phosphate transfer by the adenylic system

5 Intramolecular phosphate transfer—During the degradation of glucose or glycogen certain reactions involving phosphorus occur in which a phosphate group already present in the molecule is transferred to another position in the same molecule. For example, glycogen is broken down into a glucose phosphate compound in which the phosphate group is attached to carbon atom r of the glucose ring. This is therefore known as "glucose i phosphate" (Glucose r P). An enzyme protein, called "phosphoglucomutase" (75), can then transfer the phosphate group to carbon atom 6, the resulting substance being glucose 6 phosphate (Fig. 13). The reaction

Glucose-1 P == Glucose-6-P



is reversible, as indicated, and its necessary cofactor is the magnesium ion (75). These two phosphate glucose esters differ from each other in various chemical properties (76).

A similar intramolecular phosphate transfer occurs in the reaction

3 phosphoglyceric acid ≠ 2 phosphoglyceric acid (77)

6 Deammation —The term "deammation" refers to the removal of an NH, (amino) group, generally from amino acids Since certain amino acids form glu cose in the body and since the removal of the NH, group is the first step in such a transformation the mechanism of deammation is pertinent to the general discussion of carbohydrate metabolism.

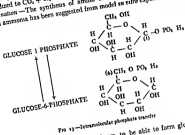
Fig 12 - Glycogen phosphorolysis

The actual loss of the NH, group from an amino acid is a spontaneous reaction not requiring an enzyme (36) However, the amino acid must first lose hydrogen before it can react with H,O to lose the NH, group (36) Hence the whole process is called an "oxidative dearmination" For example, an enzyme system known as 'amino acid oxidase, 'consisting of a protein and a coenzyme of the alloxazine group removes two hydrogen atoms from the aC atom of alanine (36, 37)

The resulting substance is known as an immo acid because of the NH or immo group. Such an acid will react with H₂O as follows

ENZYMATIC MACHINERY OF CARBOHYDRATE METABOLISM The final result is the formation of pyruvic and and ammonia (37) The NH, The lines result is the formation of pyruvic acid and animomia (37). After A13) produced may be excreted as such or transformed to urea. The pyruvic acid is

produces may be exercised as such of transformed as the H either oxidized to $CO_3 + H_3O$ or built up into glucose or glycogen. ther oxidized to CO; + 17,0 or pull up into gaucose or Eyeogen
7. Amination—The synthesis of amino acids from the corresponding Leto 7 Ammanon—Ant synthesis of annin acas from the concessioning acts and ammonia has been suggested from model in rathe experiments (78), and



one enzyme preparation has been shown to be able to form glutamate from

Although other enzymes of this kind remain to be isolated this type of reaction Although other enzymes of this kind remain to be isolated this type of reaction must be quite general for Schoenheimer has shown that following the feeding of nuss or quite general 101 octobes to experimental animals the isotopic nitrogen a labeled NII, salt (N° 18010pe) to experimental animals the isotopic nitrogen a mosero N11, sait (N° 15010pe) to experimental animals the isotopic nurgen is found in the amino groups of all the amino acids (except lysine) of their tissue is found in the animo groups of all the animo deads (except, syster) or their ussue proteins (80-81). That extensive amination must occur is also shown by the fact. proteins (so os) a mar extensive animation must occur to also shown by the fact that the corresponding lettor by droxy acids may be substituted in the diet for the that the corresponding serio or madrousy across may be substituted in the diet for the essential amino acids (82 83) Thus \ II, like CO long considered to be merely a essential annino acids 104 031 x his 3113 are consumered to so incretely a waste product to now known to be able to re-enter the metabolic cycle and func waste product is now amona to so also to texture the measure system and united that again. This must be taken into account when the urmary exerction of nitrogen is used as an index of protein catabolism (p. 127)

8 Transamination—Another type of reaction involving amino acids and related to carbohydrate metabohsm is the mutual exchange of amino and leto groups between certain a keto acids (derived from carbohydrate breakdown) and certain specific amino acids (84, 83, 86). For example

This interchange is another link between carbohydrates and protein derivatives and provides a means for the transformation of one amino acid into another. It

probably also represents a channel through which the amino acids contribute to the common metabolic pool formed by all the foodstuffs (see p 54)

9 Hydrolysis —This type of reaction is very common in the processes of digestion in the gastro-intestinal tract. Water is added to a molecule in such a way that the molecule is split into two portions, one receiving the H, the other the OH group, of the HiO (9 87). Thus sucrose, a disacchande consisting of one molecule of glucose and one of fructose, is split into its constituent hexoses by the enzyme invertiese (88). The glucosidic linkage is opened by the entry of the elements of HiO (Fig. 14).

Other examples of hydrolysis are

Lactose

Glucose

Glucose

Glucose

Glucose

However, many reactions which formerly were thought to be examples of hy drolysis have recently been shown to be phosphotolysis, e.g., glycogen breakdown (see p. x5).

THE OXIDATION OF THE HYDROGEN REMOVED FROM THE SUBSTRATE

The final products of metabolism are substances which cannot be broken down further by the tissue cells These are urea, CO, and H, Of these, urea and CO, are excreted via the kidneys and lungs respectively. The problem that remains is the final fate of the H, removed from the foodstuffs by the coenzymes (hydrogen ac ceptors). To the best of our present Lnowledge the sequence of events is as shown in Figure 15. The coenzymes are DPN, TPN, and flavin Although we are not in full possession of all the details, it may safely be assumed that the reduced pyrime nucleotides are relieved of their H, by flavin enzy mes (20, 38, 89). A final common path for H, is reached, and all of it emits as Flavin H; for an instant The scene shifts now to a series of iron-containing proteins, the cytochromes (90, 91, 92), and the "respiratory ferment" known as "cytochrome ordises" (93, 94, 95). The iron in these substances is in organic combination, in a group resembling the heme of hemoglobin (91). The iron can oscillate between the reduced and oxidized form.

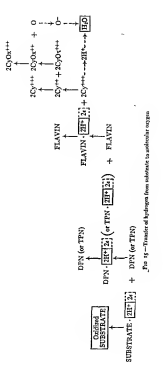
by the addition or loss of an electron. The II, of the foodstuffs, having armed at the flavin stage, reacts with the oxidized cytochrome

The electron reduces CyFe+++, while the H+ remains in the medium. The reduced cytochrome (CyFe++) reacts with cytochrome oxidase.

This serves to restore the oxidized cytochrome and to reduce the oxidise. This oxidise is unique in that it can react with molecular oxygen dissolved in the cell (63, 95).

$$OxFe^{++} + O \rightarrow OxFe^{+++} + O$$

The oxygen keeps the oxidase in its oxidized form and gains an electron. The free H^+ available from the flavin H_* then reacts with O^- to form H_*O . Thus the overall change resulting from the whole series of inversible transformations is



The series itself has been a succession of electron transfers in which every step has tended to restore the previous step to its original state

CATALYSIS BY METABOLITES

In our previous discussion of the oxidation of substrates we emphasized the role of the so called "coenzymes" as hydrogen and electron transporters. They function in this way because of their ability to be reduced and then to be reoxidized so that they may serve again Many substances of a similar nature (e.g., dyes like methylene blue) can function as electron mediators in certain in vitro biological systems under suitable conditions (96, 97, 98) These are artificially constructed pathways The cell contains certain oxidoreduction couples that can and do act like the coenzymes or the dyes (oo) For example, let us again consider the oxidation of lactate to pyruvate Diphosphopyridine nucleotide serves as the coenzyme and is reduced thereby to DPN-H. The latter is reovidized by flavin, which be comes Flavin. H. The reduced flavin may be reoxidized directly by a cytochrome, or it may be reoxidized by the couple Fumarate = Succinate The succinate, in turn, is reoxidized to fumarate by a specific enzyme and cytochrome C. The picture of events is as follows

It is, therefore, possible for a pair of metabolites to sers e as electron and hydrogen mediators in a fashion analogous to coenzymes (6, 60). This explains why, under certain conditions, a very small amount of succenate or furnarate will stimulate oxygen consumption (100, 101) The phenomenon is referred to as the "catalysis by C. dicarboxylic acids" (00, 101)

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CHAPTER III

THE INTERMEDIARY STEPS IN CARBO-HYDRATE METABOLISM

UR knowledge of the intermediary steps in carbohydrate breakdown and synthesis is by no means complete However, many lines of evidence derived from studies in vivo and in vitro in animals and in plants are con verging toward a generally accepted scheme (1, 2, 3). This scheme is outlined in Figures 16 and 17, which include the most thoroughly studied and in all probability, the most important pathways Others have been suggested and discarded from time to time. But, of these, only certain pathways for which some evidence ensist will be mentioned. It should be remembered that the present scheme is subject to revision as to detail as new data appear and that it may not apply in its entirely to all organs or tissues which utilize carbohydrates (1). One or another of the en ayme systems may be missing in a particular tissue, thus modifying the ioter mediates or the end products. The scheme, therefore, should be regarded merely as an architect s preliminary sketch, showing the general size and shape but not the final plants of the edifice to be erected.

It may be seen from Figures 16 and 17 that the orderly progression of carbohydrate breakdown can be divided conveniently into two parts (1) down to the

r from

phorylated three carbon atom units (4) At this point the first oxidative step oc

$$C_cH_{10}O_t + ATP + 2(DPN) \rightarrow 2CH_1 \cdot CO COOH + 2(ATP) + 2(DPN H_1)$$

It should be noted that one molecule of ATP was used for phosphorylation but that two molecules were formed as a result of the oxidation of phosphoglyceraide hyde and the dephosphorylation of phosphopyruvic acid respectively. This gam a ATP represents the useful energy of catabolism, as will be discussed in detail later (pp 6off) Meanwhile two molecules of DPN have been reduced, and in order to function again these must be recondized. In the presence of sufficient oxygen this is probably accomplished by a flavoprotein. When oxygen is lacking,

the pyruvic acid accepts the hydrogen of the DPN-H, and is thereby reduced to lactic acid. These two alternatives may be indicated as follows

- (1) 2(DPN · H₂) + Flavin + Cytochrome, etc + O₂ → 2(DPN) + 2H₂O
- (2) 2(DPN · H.) + 2CH. CO · COOH → 2(DPN) + 2CH. · CHDH · COOH

Thus it is clear that lactic acid is not an obligatory intermediate of carbohydrate metabolism But the breakdown of hexoses to lactic acid (glycolysis) can produce useful energy and can sustain cell functions during short periods of relative or absolute aporta

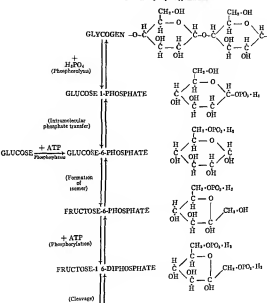
The last step above pyruvic acid, namely, phosphopyruvic to pyruvic acid, probably differs from all the others in being irreversible. It is thought that when pyruvic acid is used for carbohydrate synthesis it is first transformed to phosphoovalacetie acid, which in its turn forms phosphopyruvie acid, thus reversing catab olism by avoiding the one way step (5 6)

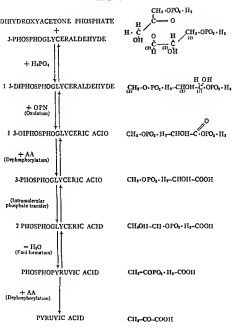
Because of the many alternative pathways which exist below pyruvic acid, the course of its breakdown to CO, and H,O is far more complex than the degradation of glucose to pyrus ate. Only the more important pathways are indicated in Figure 17 The orientation toward one or another path at a particular time will be determined by the equilibrium conditions, availability of catalysts, etc. Despite this confusing multiplicity there has emerged from the work of Szent Gyorgyi (7), Krehs (2), Barron (1, 8), Wood and Werkman (0), and Evans (10) a principal scheme of pyruvate hreakdown to CO, + H,O which is logically consistent and which helps to integrate the separate metabolisms of the three major foodstuffs

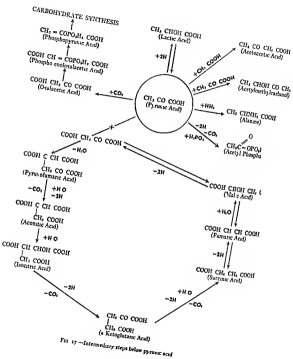
This scheme, the so-called "tricarhoxylic acid cycle," envisages the formation of a six-carbon atom acid (isocitric?) by the condensation of one molecule of pyruvate with one molecule of ovalacetate. The oxalacetate is itself formed from pyruvate hy the addition of CO, (p 34) or hy the deamination of aspartic acid The isocitrate formed goes through a cycle of oxidations and decarbox, lations un til one molecule of oxalacetate is regenerated. The latter can then start the cycle off again. It will be noted that the cycle begins with one molecule of oxalacetate and one of pyruvate and ends with one molecule of oxalacetate. In other words, in one revolution of the cycle a molecule of pyruvate has been dissimilated, and 5(II,) and 3(CO,) have been produced. The over all reaction can be written as follows

1 ozalacetate + 1 pyruvate +
$$3H_1O \rightarrow 5H_1 + 3CO$$
,
 $5H_2O$

The exact mechanism of these steps is not completely understood, but there is evidence that many of the oxidative steps involved are coupled with phosphory lation, so that eventually ATP is formed (6, 11, 12) (for significance see chan iv)







THE FINAL COMMON PATHWAY OF METABOLISM

The tricarboxylic acid cycle may assume a significance far beyond its function in carboby drate breakdown. Many amino acids may be transformed directly or indirectly into one of the constituents of the cycle. Conversely amination of the members of the cycle leads to the building of amino acids. Furthermore the recent work, of Wieland (13 14) and of Brusch (15) suggests that acetoacetic acid do net rived mostly from fatty acids may condense with oxalacetic acid to neith the same cycle. Pyruvic and oxalacetic acids and their derivatives may therefore be re garded as forming the hub of the metabolic apparatus of the cell. The cycle is probably the final common pathway for carbohydrate protein and lat as well as the locus for interconversions between the three foodstuffs (Fig. 18). With this in mind much of the older controversy as to the interconvertibility of the loodstuffs (eg. fat to carbohydrate) becomes pointless (see chaps 2n and 2nii).

ALTERNATIVE PATHWAYS

While the overwhelming mass of evidence supports the metabolic scheme out free above there are strong indications that alternative pathways may exist. For example, in certain lower animal forms (fungs and hacteria) glucose may break down without the intercession of phosphorylations (16-12). Non phosphorylative glycolysis does not seem to be spinicant in vertebrate tissues so far as they have been examined (18). On the other hand, there is indirect evidence that (under special circumstances in brain and sketelal muscle) the heroses may be completely ovidized to CO, and H,O without the intervention of the steps leading to pyrin ate formation (ing 20-21). It has been shown that complete oxidation proceeds unhampered in the presence of special militators which stop glycolysis completely. Although the alternate pathway has not been established there is some evidence to support the theory that hexose-6 phosphate may be onduced directly (22-23). Tigure 19 is a schematic representation of this hypothesis.

CRITIQUE OF METABOLIC SCHEMES

The goal of the enzyme chemist is to separate the various catalytic systems to purify them to establish their chemical properties and to study the catalyzed reactions in a homogeneous medium in wino This analytical outlook and proce dure has enriched and will continue to add to our knowledge of the metabolic mainteney of the cell in so far as the detailed properties of its parts are concerned However as in any other organized system the mere aum of the parts does not reveal the properties of the system as a whole. In the living cell which is not a bomogeneous system surface phenomena interaction between enzyme systems and other modifying influences may interfere with certain catalytic systems and promote others. For example, the rate of reparation of an intact cell is far smaller than the catalytic rate of the enzyme systems in the isolated state (8).

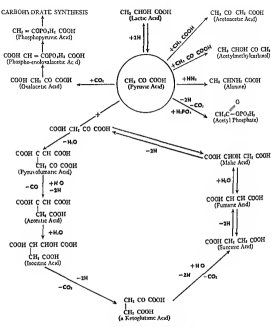


Fig. 17 -- Intermediary steps below pyruvic acid

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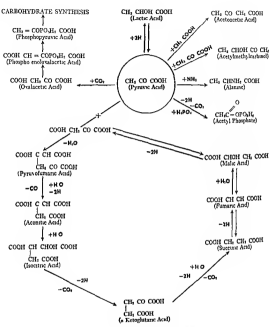


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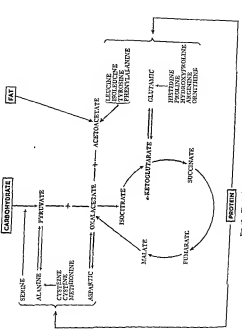


Fig. 13 -The final common pathway of metabolum

An essential characteristic of the living cell is that its metabolism is regulated Of course the rates of reactions in the cell depend upon the relative concentra tions of the activating proteins their coenzymes and the mineral elements (P Mg Fe etc) But many of the activating proteins in the carbohydrate scheme seem to depend for their activity upon sulphydryl groups (8 24) Oxidation of these groups leads to a loss of enzyme activity. It is therefore probable that the glutathione of the cell serves as a regulator of activity for many systems

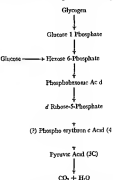


Fig. 10 -Alternate pathway for carbohydrate dissimilat on

Ever since Pasteur described the phenomenon it has been known that oxygen modifies the rate and direction of carbohydrate breakdown. In the absence of oxygen most tissues rapidly break down glycogen or glucose to lactic acid in its presence carbohydrate breakdown is slower and I ttle or no lactic acid appears The explanations of the mechanism of the Pasteur phenomenon are many and varied (25 26) In all probability there is no single mechanism for the total effect Oxygen may act by (1) removing lactic acid or its precursor (pyruvic) by oxida tion to CO and H O or by resynthesis to carbohydrate (2) maintaining some en zyme system in an inactive state by keeping (indirectly) the protein sulphur groups in the S S state (3) inhibiting the usual pathway of breakdown of carbo

CHAPTER IV

THE LIBERATION AND TRANSFER OF THE ENERGY DERIVED FROM CARBOHYDRATE BREAKDOWN

HE total energy available from the complete hreakdown of a molecule of a foodstuff to CO, and H₂O is inherent in its chemical structure. The same amount of energy would be necessary to synthesize that foodstuff from CO, and H₂O Hence, the energy can be said to reside in the chemical bonds which link the atoms to form the complex molecule. Different chemical bonds vary qualitatively and quantitatively. Some bonds are more stable than others and are therefore less reactive. A substance held together largely by such bonds is one from which the energy is less available than that from substances with unstable bonds. Different chemical bonds also vary in the amounts of energy they represent. In general, the high energy bonds tend to be the most unstable or reactive.

According to the first law of thermodynamics, no more than the total bond en ergy of a substance can be derived from its complete breakdown, regardless of the pathway or the number of intermediate steps through which this occurs. But common experience tells us that the form of the energy can be changed. For instance, the living organism can transform the original chemical energy of a foodstuff into mechanical energy (e.g., movement) Physiologists have long known that the body also produces electrical energy (e.g., nerve impulses) When the chemical or bond energy of a substance is released, it raises the temperature of the medium in which the chemical reaction takes place We speak of this as a "transformation to beat" The body temperature of animals is maintained by a multitude of such reactions There are other reactions in which the converse is true, i e, energy has to be supphed from an outside source in order to make these reactions proceed. In the laboratory we generally supply the energy in the form of beat and call such reactions "endothermic," in contrast to the "exothermic" reactions, which give off heat. In the living organism, where temperatures are very constant, the energy necessary to make some reactions proceed is applied not as heat but as chemical or bond en ergy It is therefore more precise to characterize these reactions as "endergonic" and to speak of reactions in the living organism which yield energy as being "exergonic" (1)

It will be evident that the algebraic sum of the energies of the endergonic and exergonic reactions involved in the breakdown of a foodstuff to CO, and H,O will be a positive sum of energy, equivalent to the total bond energy of the original

substance Under conditions in which this energy or any part of it has not been transmitted to objects outside the body, it finally appears and can be measured as body heat Upon this basis it has been possible to estimate total energy production (or requirements) of animals and man, under various conditions of rest and work, by measuring the total heat produced in suitable calorimeters. By simultaneously measuring the total oxygen consumption of the organism it bas also been possible to establish a caloric equivalent of the oxygen used. The estimation of the rate of metabolism from the rate of oxygen consumption is known as "indirect calorime try."

It is obvious that neither the total heat produced nor the total oxygen consumed by the body during a given period of time can give any insight into the various forms through which the original energy bas passed, nor can they indicate what bodyl functions have been served. The situation is analogous to the measurement of the heat produced by an electric light bulb made of opaque glass and of unspecified internal construction. From the total heat given off one could calculate the amount of electric current which must have been used by the bulb, and perhaps also the amount of coal which it must have taken to produce that much electrical energy. But one could not tell the amount of light present inside the bulb

SPECIFICITY OF ENERGY SOURCE

It has been customary to speak of metabolic energy as if it were an undifferentiated reservoir of power serving all cellular functions in a non specific way. How ever, recent evidence has indicated that this is not so Particular functions require particular sources of energy Indeed, they may require that the energy be derived from a specific chemical reaction. This is not surprising when one compares the situation with that which obtains with regard to internal combustion engines. If one takes a quantity of gasoline and a quantity of fuel oil of the same caloric equity alent, the former could be transformed into useful mechanical energy by a motor car but not by a Diesel powered truck, while the fuel oil would be useful in the truck and not in the car A striking example of the specificity of fuel in the living organism is the essential nature of plucose for the activity of the central nervous system When isolated brain tissue is studied in vitro by the Warburg technique. it can readily be demonstrated that its oxygen consumption (energy production) can be as well maintained at the expense of pyruvate or succinate as by the use of glucose (2, 3, 4) Nevertheless, in the intact living animal the brain evidences senous functional difficulty as soon as the blood sugar level falls below about 40 mg per cent Apparently, the normal irritability of the central nervous system depends upon chemical energy derived from glucose. This function cannot be main tained at the expense of the energy derivable from lower intermediary substances (5, 6, 7)

THE ENERGY TRANSFER FUNCTION OF PHOSPHATE GROUPS

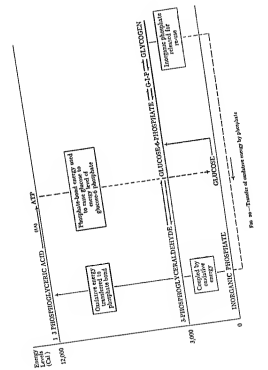
It is now known that the various phosphorylations which occur throughout the dissimilation of carhohydrate are the means by which the energy liberated from oxidative steps is prevented from being dissipated as heat and is held or built up for use in endergonic reactions (8, 9). Different phosphorylations carry different amounts of energy and are, therefore, suitable for motivating different kinds of endergonic reactions (9). According to the amount of energy transferred, we speak of high energy or of low-energy phosphate compounds or bonds. Inorganic phosphate is of course, at the lowest energy level. The high energy phosphate bonds (10,000-12,000 cal/mole) are present in such compounds as adenosine triphosphate (ATP), creatine phosphate, acetyl phosphate, phosphopyriuse, etc. As an example of how a high energy phosphate bond performs its function, let us con sider the manner in which glucose is transformed into glycogen, a carbohydrate of higher potential energy than its precursor. A superficial representation of the chemical steps between elucose and elucose much be written as follows

Glucose -> Glucose-6 phosphate == Glucose-1 phosphate == Glycogen

From an energetic standpoint this reaction by itself is impossible, since it requires the addition of energy to raise glucose to the energy level of glycogen, and there is no indication whence this energy is derived. These reactions can be made to proceed in vitro by adding certain protein enzymes and ATP (10, 11, 12). The energy which drives the reactions is derived from the high-energy phosphate bonds in the ATP. The latter loses its lablic phosphates, becoming a denivel acid in the process.

Since the amount of ATP present in living cells is limited, the more complete story of the series of reactions in the living organism must include the manner in which adenylic acid is rephosphorylated to ATP. This may occur in more than one way, but an important means is through the energy liberated by the oxidation of 3 phosphoglyceraldehyde to 1 in 3 phosphoglyceraldehyde to 4 in a central color of the addenyde to the acid is incorporated in a high energy phosphate hond in the acid. In a sense, therefore, we may say that the oxidative energy has raised the morganic phosphate involved in the reaction to a higher energy level (6). The motivating power of the chain of events having thus been applied, the cycle proceeds in the manner graphically illustrated in Figure 20. It may be seen that the ultimate use of the original oxidative energy, applied through ATP, is to raise the lower energy foodstuff (glucose) to the higher-energy storage product (glycogen). At the latter point the phosphate group involved in the sense of reactions is divorced from the substrate and may re-enter the cycle at the beginning.

The raising of glucose to the energy level of glycogen is only one of the functions which ATP performs Indeed, the reversible systems AA == ADP == ATP seem to



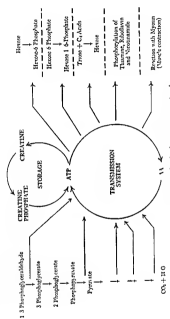


Fig. 21 -Central position of the adenylic system in energy transfer

be the central mechanisms for energy transfer from exergonic to endergonic reactions in carbohydrate metabolism. Figure 21 summarizes their relationships to all the known energy cycles.

There has been considerable doubt as to the place of the Creatine == creatine bonds, some authors have ascribed to creatine phosphate a role similar to that indicated for ATP. It now seems more likely that the latter is not the case but that creatine phosphate acts as an emergencystore ofling be energy phosphate bonds. This store is built up at times when the AA == ATP systems are producing an excess of energy over the requirements of the moment and is broken down when the ATP mechanisms cannot supply energy as tapidly as is required. Thus, creatine phosphate stands in the same relationship to the storage of energy as glycogen stands in relation to the storage of carbohydrate substrate.

Finally, it should be noted that the transference of energy by means of phos phate bonds accounts for the ready reversibility of most of the reactions of carbo hydrate metabolism (8, 9, 14). This is because the energy which is yielded by the substrate remains "attached" to the product of the reaction and is therefore not lost from the system For example, the hydrolytic splitting of glycogen by amplopably produces glucose and hiberates energy as heat. The analogous phosphorlytic cleavage of glycogen in the body (see Fig. 12, p. 38) produces glucose I phosphate, with the energy retained in the phosphate bond. Hence, no outside energy is necessary to reverse the process (8, 15).

Regarded as a whole, the pattern of energy interchange in carbohydrate metab olism is by no means as complicated as a consideration of the details might lead one to believe The general principle may be compared to that employed in the mining and use of coal Figure 22 is a diagrammatic representation of the anal ony, in which various features are labeled with their metabolic counterparts. The essential features are the investment of a certain amount of energy to procure large amounts of an energy substance (coal in the mine shaft, or glucose in the body), the raising of the energy substance to a higher energy level (the coal pile on the surface, or glycogen in the body), the conversion of the energy substance into another form of energy (running the electric general it from a steam engine fired by coal, or phosphorylation in the body), the use of the more convenient form of energy for the transfer of power to places where it can be used for special purposes (use of electric power for communication, transportation, etc., or the use of phos phorylative energy for muscle contraction [8], nerve conduction [16], intestinal absorption [17] renal reabsorption [18], calcification [19], sperm motility [20], etc), and, finally, the use of some of the energy derived from the energy substance to obtain more of the energy substance (use of some of the electrical energy made from the coal for the purpose of mining more coal, or the phosphorylation of glu cose in the body)

Since we do not, as yet, possess a detailed knowledge of all or most phosphateenergy transfer reactions, the efficiency of this mechanism can be judged only anproximately It has been shown that, during the complete dissimilation of I mol of glucose to CO, and H,O, from twelve to twenty-four high-energy phosphate bonds are formed (21, 22, 23, 24). The energy content of these phosphate bonds is. therefore. 144,000-288,000 cal Since 1 mol. of glucose going to CO, and H₂O yields 673,000 cal, the energy transferred by means of phosphate bonds represents about 21-42 per cent of the total It is interesting to compare these figures with that of the efficiency of muscular work, which is generally considered to be about 30 per cent

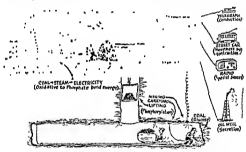


Fig 22 -Analogy to the liberation, transfer, and utilization of metabolic energy

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CHAPTER V

THE USE OF ENERGY FOR MUSCULAR CONTRACTION

IN THE previous chapters we clarified our concepts as to the nature of the energy derived from carbohydrate and the manner in which this energy is transformed and made available for the various uses to which it is put The actual results of the expenditure of useful energy in the body may be observed in terms of muscular contraction, glandular secretion, nervous activity, etc. It remains to consider in detail how the very real but invisible energy of the foodstuff is translated into tangible physiological performance. Muscular contraction will serve as the best example for this purpose. This is partly because more is known about this function than about any other and also because it is quantitatively the most important energy outlet.

Skeletal or voluntary muscle comprises approximately 50 per cent of the body weight It consists of 75-50 per cent H₁O and 20-25 per cent solids The dry unish of the muscle is partitioned as follows (omitting lipiods and mingrals)

75-80 per cent proteins

2 5-5 o per cent glycogen 2 0-3 o per cent creatine phosphate and free creatine

1 0-1 5 per cent adenosine phosphates

I o per cent other phosphorylated products of car bohydrate metabolism

It may be seen that protein is the chief structural component of this tissue. But it must be remembered (as pointed out in chap ii) that most, if not all, of the proteins of the living cell function as enzymes as well as structural elements Next to protein in quantitative importance are the two storage products, glycogen (the fuel reserve) and creatine phosphate (the more readily available energy reserve). Adenyla caid and its phosphorylated forms, which constitute the active phosphorylating system of the muscle, represent a small but significant fraction of its bulk. The remainder of the muscle is composed of a number of intermediate metabolities which are caught in transit.

THE PHYSICAL NATURE OF MUSCLE CONTRACTION

The contractile element responsible for the shortening and elongation by which muscle performs its physiological function is myosin, one of its proteins (1, 2) Myosin is present in the form of elongated, threadlike structures called 'muscle fihrils." These are microscopic in size A bundle of fibrils, composed of large num bers in parallel formation, constitutes a muscle fiber. The gross structure of a muscle is composed of aggregates of fibers. The myosin of the muscle fibrils represents approximately half of the total muscle protein.

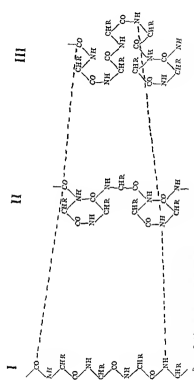
Both in its shape and its elastic properties the myosin fibril resembles a rubber band (3, 4). It is not unique in this for keratin and wool are proteins of the same type. But myosin differs from these other proteins in having an internal mechanism by which it is stretched. The contraction of a fibril is due to the release of this mechanism and to the fibril's recoil to a neutral position. X ray diffraction studies have indicated that the internal configuration of the myosin molecule, in its stretched and collapsed states, changes as shown in Figure 21 (4).

It will be noted that a relatively new and unorthodox conception of muscle states has been introduced in the preceding paragraph. It has been customary to speak of a resting muscle as "relaxed" and of a working muscle as "contracted". As these terms imply, it was formerly thought that the energy expended in work was applied in bringing about the shortening or contraction of muscle, while re laxation was merely the result of the cessation of the expenditure of contractile energy. The newer evidence, that the resting muscle resembles a stretched elastic hand, necessarily reverses the locus of application of energy. The external force exerted by the contracting muscle is a result of the recoil of its stretched fibrils, while the metabolic energy is applied to return the collapsed elastic members back to their orienal state of stretch.

THE CHEMICAL EVENTS ACCOMPANYING MUSCLE CONTRACTION

The first chemical changes to be related to the change in the physical state of the muscle during contractors were the breakdown of glycogen and the appearance of lactic acid (5, 6). Lundsgaard's demonstration that contraction of muscle was possible in the presence of iodocctate, which prevented lactic acid formation, forced the abandonment of this hypothesis. He further demonstrated a parallelism between the breakdown of creatine phosphate and the energy liberated by the iodoccetate treated muscle. This led to the hypothesis that the immediate source of energy for muscular contraction was the hreakdown of creatine phosphate, while the glycolytic process served to resynthesize the creatine phosphate from its split products (7, 8, 9).

The current conception of the means by which metabolic energy is applied to the muscle fibrils was initiated by the work of Lohman, who showed that adenosine triphosphate (ATP) was necessary both for glycolysis in muscle and for the syn thesis of creatine phosphate (io, ii) This was followed by Parnas' demonstration that the hreakdown of creatine phosphate merely served to supply phosphate for the conversion of adenylic acid to ATP, without the hieration of energy, while the subsequent hreakdown of the ATP actually supplied the energy for contraction



n a r-Septembel and contracted forms of the myon modecule I artificantly verechood to time of extremely by mechanical means II as the myonomelous is thosely to scart to retained state meno III as at a state from the contracted state meno III as at a state from the Representative state which chan each state when R represents the surrous stand side.

and the phosphate for glycolysis (12–13). The glycolytic reactions in turn provided the energy for the resynthesis of both creatine phosphate and ATP

It may be seen that as our knowledge of the subject has developed the break down of glycogen to lactic acid has been gradually relegated to a secondary process with a restorative function. As a matter of fact the most recent evidence indicates that under ordinary physiological conditions glycogen breaks down without the appearance of lactic acid at all (p. 49). When the rate of oxygen supply to the muscle is adequate for the rate of glycogen breakdown pyrivic acid is oxidized completely and none of it is reduced to lactic acid. Under these conditions oxida tive steps above and below pyrivine acid supply energy for the rephisphorylation of ATP and thus maintain the metabolic cycle in the absence of lactic acid. It is only when the oxygen supply is inadequate (as it was in most of the experiments of the earlier investigators) that lactic acid appears. This occurs because pyrivic acid partially substitutes for oxygen by becoming the hydrogen acceptor from reduced DPN and in so doing is itself freduced to lactic acid.

In a sense therefore the formation of lactic acid by muscle is merely an emergency mechanism enabling muscular contractions to occur for a short time despite a lack of oxygen. This may be useful at the beginning of sudden or severe muscular work to tide the muscle over a period of circulatory adjustment that is while the blood supply is changing from the slow rate adequate during rest to the more rapid rate necessitated by the exertion (14). It also enables the muscle to exert a relatively tremendous effort for a short space of time at a rate with which the maximal rate of oxygen supply could never cope. The lactic acid which accumulates during such an effort is reoxidized to pyruvic acid when the exertion is over. This process may be regarded as the repayment during comparative lessure of an energy delt contracted under stress.

Figure 24 graphically illustrates the development of our concepts concerning the sequence of chemical events which occur during muscular contraction

Although it is out of place here to attempt an analysis of conflicting data in respect to the chemistry of muscular contraction (as it occurs in riso), it should be
pointed out that the work of Sacks (is) and of others (io) indicates that the
schence as given in Figure 24 may have to be modified to account for the sequence
of chemical events in the living intact muscle

THE CONNECTION BETWEEN THE PHYSICAL AND CHEMICAL EVENTS IN MUSCLE CONTRACTION

Thus far we have merely described the chemical events which occur coinci dentally with muscular contraction. It remained for Engelhardt (17, 18) to demon strate the direct causal link between the chemical reactions and the change in the phys cal state of the myosin. In so doing, he confirmed the dominant position of ATP in the chemical processes as well as the previously described physical nature.



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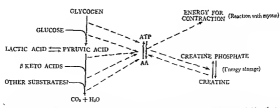


Fig. 24—Descionment of concepts of the chemistry of muscular contraction I. Hopkins-Meyerhof hypothesis, Lundsgaard modification, III. current scheme indicating the scondary role of lattic and, the critical positions adought system, the energy storage function of cerainer phosphate, and the use of the energy in ATP by myonal

of contraction (i.e., the recoil of a stretched fiber) By injecting a thin stream of a purified myosin preparation into water, Engelhardt (18) was able to make threads of myosin analogous to muscle fibrils and possessing similar elastic properties. When suitably weighted and suspended in water, these myosin threads were not affected by the presence of the various mineral and organic substances normally found in mammalian muscle. But the addition of ATP to the water was followed by a definite increase in the length of the threads which could be reversed by flushing away the ATP.

Szent Gyorgy and his co workers (19) confirmed Engelhardt's work and extended it into a more complete analogy of 18 time muscular contraction. They found that a purer preparation of myosin than that used by Engelhardt would not form threads when injected into water. But when another muscle protein (which they named 'actine') was added to the myosin, the compound behaved his Engelhardt's preparation. They named this complex 'actomyosin' and found that threads formed from it could be made to extend or contract at will by varying the proportions of ATP potassium, and magnesium added to the water in which they were suspended

The extremely simple conditions of Engelhardts and Szent Gyorgy's experiments leave no doubt that ATP is the prime agent responsible for the Stretching of myosin fibrils that is preparatory to muscle contraction. The peculiar appropriate ness of ATP for this purpose hes in the fact that it had previously heen shown that myosin is the enzyme which splits ATP — ADP + P, 6,0, 2;). For the time he ing, we may therefore accept the current scheme shown in Figure 24 as representing the cycle of events by which metabolic energy derived from the utilization of carbohydrate is transferred by ATP and applied to the contractile elements of the muscle. The train of reactions is such that both the original abusined that of the muscle and the original amount of ATP are restored subsequent to contraction.

It is evident from our present conception that any metabolic intermediate which can supply the energy necessary to restore AA to ATP can serve as a fuel of muscular exercise. This applies to a and \$\beta\$ Letoacids derived from protein and fat as well as to carbobydrate derivatives (see Fig. 18, p. 54).

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PART II INTRODUCTORY PHYSIOLOGICAL CONSIDERATIONS



CHAPTER VI

NATURE AND OCCURRENCE IN THE TISSUES OF MATERIALS IMPORTANT TO CARBO HYDRATE METAROLISM

In THE previous chapters we discussed the ultimate use of carbobydrate by the effector organs and the mainter in which the chemical energy of the food stuff is liberated and applied to physologic functions. It will readily be appreciated that this knowledge however fundamental and important, is only a small part of the larger body of information with which it must be integrated in order to understand carbohydrate metabolism in the living organism. As opposed to chemical reactions in the laboratory, an essential characteristic of metabolic functions in vious that they are finely regulated processes adjusted in each organ and tissue to the needs of the body as a whole. It is with the complex series of actions and interactions between tissues and organs subject to intrinsic endocrine and nervous regulation that we must now deal. But before beginning our account, it will be useful to describe in some detail the nature and occurrence in the tissues of various substances which are important to carbohydrate metabolism—substances which have been briefly mentioned in the preceding chapters and which we shall meet again in subsequent chapters.

CLUCOSE

Glucose is the chief and for practical purposes the only, transport form of carbohydrate Carbohydrates enter the blood from the gastro intestinal tract largely as glucose in the post absorptive state, glucose is the carbohydrate which the liver supplies to all the other tissues of the body. For these reasons the level of glucose in the blood is normally higher than in any other tissue or fluid of the body.

The average normal level of glucose in the blood does not vary appreciably with the species of animal In most mammals it is very similar, ranging from 60 to 80 mg per 100 cc of whole blood It has been customary to express these amounts as '60-80 mg per cent' Strictly speaking, this is incorrect, for whole blood is not homogeneous nor is tof the same specific gravity as water Nevertheless, with the reservations noted we shall make use of this shorthand designation of concentration for the sake of convenience

The blood sugar levels reported by different observers depend to a certain extent, upon the methods employed for chemical analysis. Glucose is an aldohexose

(see Fig 25) in which the aldehyde group on the first carbon atom acts as a reducting agent. Hence, the most practical and most commonly used chemical methods for determining glucose are procedures in which a metallic ion in the oxidized state (usually copper) is feduced by the sugar. Such methods were devised by Bertrand (i), Folin (a), Hagedorn (3), Somogyn (4) and many others (5, 6). They differ from each other chiefly as regards the means by which reducing substances other than glucose are removed from the reaction. To the extent that these means differ in efficiency, there are differences in blood sugar values reported from various laboratories. For example, the range of normal values quoted for mammals is obtained by the Somogyn modification of the Shaffer Hartman method. When the Folin Wu method is used, a range of from 80 to 120 mg per cent is obtained Somogyi has shown (4, 7) that his method of precipitation removes virtually all the non carbohydrate reducing substances (chiefly glutatione), hence, the results

obtained by using his method are sometimes referred to as values for 'true' blood surar

When the level of sugar in a sample of whole blood is 100 mg per cent, the concentration of sugar in the plasma of the same blood is about 115 mg per cent (8, 9). This difference is due to the fact that the sugar is not equally distributed between the blood plasma and the red blood cells. (There is an equal distribution of glucose between the blood plasma and the water phase of the red blood cells [8, 20]. The precise difference between the whole blood sugar and the plasma sugar in a given instance will depend upon whether or not the normal number of red blood cells per unit volume of blood is present.

Because the peripheral tissues are constantly removing sugar from the blood samples of arternal or capillary blood will show a level of sugar a few milligrams per cent higher than that of simultaneously drawn samples of venous blood (11, 12). This so-called A-V difference varies with the ensting rate of utilization of sugar and also depends upon the rate of blood flow through the tissues at the time of sampling (13, 14). It is obvious that, if the rate of sugar utilization were con

stant, a doubling of the rate of blood flow would result in a diminution of the A-V difference to half its former value Neglect of this simple consideration has given rise to some confusion in the literature (14, 15)

Table y lasts the range of sugar values reported an various fluids and secretions of the body Being a crystalloid of small molecular weight, glucose diffuses readily out of the blood stream into all other body fluids. Tessues like liver or skeletal muscle are composed of at least two phases, namely, the tissue cells and the fluid filling the interstices between them (extracellular fluid). The sugar in the blood plasma would rapidly equilibrate with the sugar in the extracellular fluid were it not for the constant withdrawal of sugar from the latter by the cells. The actual level of glucose in the extracellular fluid is therefore a few milligrams lower than that in the blood plasma. But analysis of normal whole muscle for its glucose con the cartacellular fluid of the proper presentions to prevent glucolysis suitally yields a range of

TABLE 7
GLUCOSE CONTENT OF BODILY FLUIDS

Flu d	(Mg per Cent)
Whole blood	60- 90
Blood plasma	70-110
Lymph	70-110
Cerebrospinal fluid	40- 70

values between 30 and 60 mg per cent This, of course means that the cells then selves contain much less free sugar than does the extracellular fluid. An estimate of the amount of glucose actually present within the tissue cells may be made by determining the amount of extracellular fluid and calculating the intracellular sugar from the sugar content of the whole tissue.

Normal unne contains a small amount of glucose An average adult human excretes from $\frac{1}{2}$ to $\frac{1}{2}$ gm in the approximately $x_1, x_0 = c$ of urne excreted in 24 hours (16 17). In chinical medicine such urne is termed "sugar free," because the routine methods for the qualitative detection of sugar are not sufficiently sensitive to undicate its presence in this concentration. That the concentration of glucose in normal urne is far below that occurring in other body fluids is not because the membranes of the kidney are less permeable to sugar. The kidney glomerulus actually passes a filtrate containing glucose in the same concentration as is present in blood plasma (18). But this filtrate is then subject to the action of the cells of the kidney tubules which reabsorb most of the sugar in it (19, 20).

The process by which the kidney tubules reabsorh glucose depends upon phosphorylating mechanisms (21, 22) Inhibition of the latter by the glucoside phlor humin prevents the reabsorption of the sugar and results in so-called "phlorhizin diabetes" (33, 24) Abnormal amounts of sugar also appear in the urine whenever the blood sugar level is raised to such heights that the amount of glucose filtering

through the glomeruh exceeds the phosphorylative capacity of the tubules. The critical level at which this begins to occur is usually about 180 mg per cent and soften referred to as the kidney threshold for glucose (20 25)

GLYCOGEN

Glycogen in the animal body is similar in form and function to the starch in plants. It is a polymer consisting of many glucose molecules joined to each other in the manner indicated in Figure 11 (p. 37). The CO-C-linkage between adjoining glucose molecules is known as the glucosidic linkage. It is here that the glycogen molecule splits with the introduction of a phosphate group (see p. 35). The distribution and particular significance of the glycogen in various tissues is discussed in a number of places throughout this book (see p. 9) and will not be repeated here.

Glycogen when isolated in the laboratory is a stable compound. But in the presence of the tissue enzyme systems it breaks down very easily. For this reason the glycogen content of a dead tissue gives no indication of its content during life and accuracy of estimation is not assured even when tissue is removed from the living organism. This is especially true when any degree of anoxia is allowed to occur while the tissue is being removed for analysis or in the case of muscle when twitthing of the muscle fibers is induced by careless handling. A probable reason for the susceptibility of glycogen to anoxia is that the active form of glycogen phosphorylase (see p. 36) contains. SH (reducing) groups. Hence any degree of oxygen lack would tend to keep the enzyme in its reduced form and would there fore favor the phosphorylation and breakdown of glycogen. Another reason may be the rapid appearance of inorganic phosphate during oxygen lack. This favors glycogenolysis by shifting the equilibrium of the following equation to the right

Glycogen + P. = Glucose-1 phosphate (26)

The standard method for glycogen estimation in tissues depends upon the fact (discovered by Claude Bernard and put to practical use by Pffuger) that het concentrated potassium hydroxide destroys all carbohydrates except glycogen As described by Good Kramer and Somogyi (27) the method is accurate and relatively simple once the tissue is dissolved in the alkali. The difficulty consists in removing and transferring the hving tissue into the alkali before any 8 g inficant amount of glycogen disappears. Fairly good and consistent results may be obtained by aneitherizing the animal with an aneitheric (such as amytal or pento-barbital) which does not itself tend to break down glycogen. The tissue to be student and the such as the such

on

taining hot alkali But it cterm qu by men e content of a tissue is most nearly determinable is the following. The animal is anesthe

tized, and the tissue prepared as above The tissue is then frozen in situ by the use of liquid air or crushed CO, ice It is removed and weighed in the frozen state and immersed in the hot alkali

LACTIC ACID

When the body is at rest and in the post absorptive state, the lactic acid content of the blood ranges between 10 and 20 mg per cent (28, 29). The lactic acid content of other tissues is in equilibrium with that of the blood plasma, for lactic acid is freely diffusible across cell membranes (30). Under these circumstances the small amount of lactic acid which is present probably arises from a few special tissues, such as the red blood cells the intestinal mucous membrane, and the ret ina, etc. Adult mammalian erythrocytes do not possess the enzymatic machinery for the use of oxygen but readily produce lactic acid from blood glucoses (31, 32). The cells of the intestinal mucosa (33) and of the retina (34) have a high aerobic glycolysis (see p. 55), that is they differ from most tissue cells, in which an ade quate oxygen supply inhibits lactic acid production (Pasteur effect [p. 55])

In most tissues of the body lactic acid is not a necessary intermediate of carbo bydrate metabolism. It is formed by the reduction of pyravic acid only when the modature removal of the latter is relatively or absolutely deficient (p. 49). A relative oxygen lack may occur during strenuous physical exercise, when the rate of oxygen supply to the muscles is temporarily madequate in comparison with the rate of glycogenolysis (30), whereupon the lactic acid in the muscles increases and diffuses out into the blood. Certain organs particularly the liver (35, 36) but also the heart (37, 38) will then remove the excess lactic acid from the blood and reconduct it to privarye acid.

An absolute lack of oxygen, leading to high lactic acid levels even when the body is at rest may result from pulmonary (39) or cardiovascular (40) diseases which interfere with the oxygenation of the blood or tissues, respectively A similar end result may be caused by liver disease (41), when the impairment of the oxida tive systems in this organ prevent it from utilizing the oxygen available in the blood for the removal and oxidation of the blood fact the removal and oxidation of the blood fact is emoval and oxidation of the shood fact is emoval.

The importance of anoua in lactic acid formation necessitates the same precautions as for glycogen (p. 78) when sampling tissues for chemical analysis. The addition of sodium fluoride to blood prevents further glycolysis (4.2) Lactic acid is usually estimated by the method of Friedemann (43) or by that of Miller and Muntz (44) The latter method was modified and adapted to tissue analysis by Barker and Summerson (45)

PYRUVIC ACID

Since pyruvic acid is one of the most reactive metabolic intermediates (see p 52), it is not surprising that the amounts of pyruvic acid normally found in the blood and other tissues do not exceed 10 mg per cent (46, 47). The level rises

somewhat with the increased hreakdown of carbohydrate accompanying muscular work (48) or following carbohydrate administration (46, 49) The pyruvic acd content of hlood and tissues also increases during thamme deficiency (50, 51), for many of the reactions which dispose of pyruvic acid require thiamine diphosphate as a coenzyme. This fact has been used as an aid in the diagnosis of this avita minosis (49, 51).

It should be noted that, despite the fact that pyruvic acid is hy far one of the most important substances in intermediary metaholism, its normal concentration in blood and tissues is only about one tenth to one twentieth that of lactic acid. This is because of the many mechanisms available for pyruvate removal (p. 53), while lactic acid disposalishmited to one reaction—its oudstion to pyruvate. This il lustrates the general rule that the concentration of a substance in blood and tissues is not necessarily an indication of its importance in the metaholic scheme. As we shall see presently, some metaholic intermediates are never present in detectable amounts unless special methods are employed to stop the metaholic reactions at that stage.

The method commonly used for pyruvate estimation is that of Lu (52), or the subsequent modifications of this method (53, 54)

PHOSPHATE COMPOUNDS

We have already discussed the predominant role of compounds of phosphone acid in carbohydrate assimilation and dissimilation (p. 60). The phosphate deriv atives group themselves into three classes imorganic phosphate, phosphorylated intermediates, and phosphate transfer substances.

Inorganic phosphate (P.) —The P. in the body is largely derived from the inorganic phosphates present in foods Under certain circumstances the P. of the blood may be increased by the mobilization of Ca₂(PO₂), from the hones The P. of blood and soft tissues may also rise as the result of an increased breakdown of organic phosphate compounds owing to anoxia or the interruption of the activity of certain enzyme systems. Hence, the sampling of tissues for the correct estimation of P., as well as of the other phosphate derivatives, involves the same precautions as for glycogen (p. 78). With more careful handling of tissues, lower P. values have been reported (55). Table 8 summarizes the most reliable observations as to the levels of P., and other important phosphate compounds in various bodily tissues.

Phosphorylated intermediates —The only phosphorylated intermediates of car bohydrate metabolism which are normally present in the tissues in detectable quantities are (a) hexose-6 phosphate, (b) monophosphoglycenc acid, and (c) diphosphoglycenc acid (in red blood cells only) Table 8 lists the levels which have been reported The other known phosphorylated intermediates, such as glucose I phosphate, herose diphosphate, etc., are metabolized as rapidly as they are pro-

duced and therefore are not found except when steps have been taken to interfere with their disposal (42, 56)

Photphate transfer substance:—This group consists of (a) adenosine diphosphate, (b) adenosine triphosphate, and (c) creatine phosphate. The levels normally found in tissues appear in Table 8 The adenosine polyphosphates are present in

TABLE 8

DISTRIBUTION OF PHOSPHATE COMPOUNDS IN VARIOUS
TISSUES OF MAN, RAT, RABBIT, AND DOG

Skeletal

Tissue	Inorgan 10 Phos- phate (Pa)	Crest se Phos- phate (CrP)	a ne Dy and Ter- phos- phates (ADP and ATP)	Herose- 6-Phos- phate (HMP)	Phespho- alycerate (PGly)	D phas- phogye- grate (d PGly)	Total Acid- soluble Phos- phate (P Total)	Refer- ences
muscle muscle	15-25 13-29 18 7-9 3 5	50-70 5-13 0 9-11	30-40 18-28 15-25 10-19 10-20	8 ts 14 4-6	40-50	30-50	150-200 80-100 90-100 70 50-80	(58,62) (64) (60) (59 66) (61)

TABLE 9
PROPERTIES OF THE VARIOUS ORGANIC PHOSPHATE COMPOUNDS
(Robison and MacParlane [71])

Frocedure	Cres t ne Phor- phate	ATP and ADP	Gha cose- s Phos- phate	Glu 605e- 6-Phos- phate	Fruc- tose- 6-Phos- phute	Hexase D phos- phate	Triose Phos- phate	Phos- pho- glyc- erate	Phos- pho- pyru vate
I Percentage of hydrol ysis in molybdate at 25° C for 30 minutes	100	0	0		0	۰	۰	۰	٠
II Percentage of hydrol yss in N HCl at roo* C In 7 minutes In 30 minutes In 60 minutes In 180 minutes		100	100	3 9	24 45 84	32 59 72 94	46 92 100	1 2 6	93
III Percentage of hydrol ys s in N NaOH at 20°C in 15 minutes						۰	100		۰
IV Reducing power per roo mg of the free ester, compared to glucose as roo	1		۰	\$5	ss	21	50	o	

all tissues of the body to a greater or lesser extent (57, 58 59 60 61) However creatine phosphate secms to be limited to contractile and conducting tissues—1 c, striated smooth, and cardiac muscle, neurones, and nerve fibers (62 63 64 65 66) It has also been found in spermatozoa (67) There is no creatine phosphate in blood or liver (60 67)

Analytical methods —The methods for the estimation of the various phosphate compounds are based upon separation of the desired compounds from each other by the differential solubility of their barium salts (68, 69) and the varying conditions under which the inorganic phosphate portion can be split off the particular organic substances (70 71, 72) Table 9 outlines the principles underlying the various determinations A reliable method for the estimation of P, must be used of course. In all such procedures (87 a 7).

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CHAPTER VII

SITE OF ORIGIN OF BLOOD SUGAR

IT IS well established that in the fasting animal the liver is virtually the sole source of the blood sugar (i. 2.3). There is some recent evidence that the kidney may contribute sugar to the blood but in amounts that are hardly ignificant in relation to the total carbohydrate requirements of the normal intact mimal (4.5). The other tissues of the body continually require and use the blood ugar for the maintenance of their metabolism and functions. Since the blood ugar level is well maintained throughout long periods of fasting it is evident that he sugar which the liver secretes into the blood under these conditions must be lerived from stored carbohydrate or non carbohydrate precursors. It has been ineffy indicated in the previous chapters that the storage form of carbohydrate procursors are protein and fat. The present and following chapters will consider the evidence for these interconversions in one detail.

The brillant pioneer work of Claude Bernard was the first to indicate the predominant role of the liver in supplying blood sugar and to demonstrate the existence of liver glycogen. His early reports claimed that in fasting animals or those fed on meat the blood entering the liver through the portal vein contained no sugar (6). Repetition of these experiments by some of bis contemporanes led to disagreement and controversy for they found sugar in the portal vein blood As it turned out the reasons for these differences lay in the then inadequate knowl edge concerning the proper handling of blood samples and the crude methods for sugar analysis. Bernard and his contemporanes eventually agreed that while sugar was constantly present in the portal blood. there was always more sugar in the blood leaving the liver (7).

Claude Bernard also demonstrated that a liver flushed free of sugar by perfusion with cold water acquired a high sugar content after a few bours in the labora tory. He recognized the starchlike nature of the precursor of this sugar and called it glycogen. He confirmed Chauveau in the finding that the sugar of arterial blood throughout the body was higher than that of venous blood. On the hasis of these essential facts and a number of other observations. Bernard arrived at the following conception which is as valid today as when he enuncated it.

In the liver sugar is produced although a little is also destroyed in that organ in the muscles sugar is destroyed. Destruction of sugar probably occurs throughout the organism in all the

THE LACTIC ACD, CYCLE

The evidence which has been cited also shows that, once sugar has entered th peripheral tissues, even though it is stored rather than used, it cannot re ente the blood as glucose This, of course, is in accord with what is known of the en zvme systems in skeletal muscle (p 34) However, under special circumstances significant amounts of carbohydrate can leave the muscle in altered form, as when lactic acid accumulates in the muscle and diffuses into the blood stream. This oc curs during a relative or absolute deficiency in the oxygen supply to the muscle (see p 49) At such times the lactic acid may be carried to the liver and converted into hepatic glycogen, and thus eventually reappear as blood sugar. This so-called "lactic acid cycle" has been investigated and elaborated by Geiger (25, 26, 27), Himwich (28), Cori (29), and others But it is fair to say that, while it constitutes a possible indirect source for some blood sugar during abnormal or emergency con ditions, it is of little or no significance as regards the blood sugar supply under normal conditions

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An entering wedge into the solution of the problem was made by Hershey and Soskin (42, 43), who showed that it was not the digestive-enzyme activity of the administered pancreas that was essential for the relief of the fatty liver and the accompanying syndrome of "liver failure," as had previously been supposed. They demonstrated the same effects by feeding a preparation of egg yolk lection Further work by Best, Hershey, and Huntsman (44) revealed that it was the choline constituent of the lecitini molecule that exerted all the physiological activity Since then, the literature on choline and other substances with similar activity ("lipotropie" factors) has grown enormously (45), and a complete review of this subject would take us far afield. What is pertinent to the present discussion is the observation of Ralli et al. (40) that the lipotropic activity of raw pancreas was greater than could be accounted for by its lection or choine content.

In 1936 Dragstedt and his associates (47, 48) began an important series of in vestigations by preparing an active pancreatic extract which, despite its low choline content, was a very effective photropic agent in the depancreatized dog They named the active principle "hipocaic" and tentatively considered it to be a hor mone, because occlusion of the external pancreatic ducts of normal dogs did not result in any evidences of the lack of the hipotropic substance. The hormonal nature of lipocaic has been challenged by the laboratories of Chaikoff (49, 59, 57) and of Ralli (46, 52), which have reported (6) that, in their hands, ligation of the pancreatic ducts does produce a fatty-liver and (6) that the oral administration of the external secretion of the pancreas (pancreatic juice) yields as great a lipotropic effect as the feeding of raw pancreas. These contradictory results and conclusions have not yet heen resolved. What concerns us for the moment, however, is the area of agreement (53, 54), i.e., that the pancreas secretes, whether internally or externally, a lipotropic agent other than, or in addition to, choline

The subject has been complicated by the use, by various investigators, of ani mals other than the dog and methods other than pancreatectomy. In a comprehen sive review of the literature on lipotropic factors McHenry and Patterson (45) reached conclusions which may be summarized as follows.

- 1 There are different kinds of fatty livers, depending upon how they are produced and differing in the chemical composition of the liver lipids (see Table 10)
- 2 When the fatty liver contains a high percentage of neutral fat, choline is an effective lipotropic agent
- 3 When the fatty liver contains a considerable percentage of cholesterol, lipo-

nicious anemia factor

Wherever the future work on hpocase and other hpotropic substances may lead, it is clear that, in dealing with the departreatized dog, one must provide hpo-

tropic agents adequate in kind and amount to prevent fatty infiltration and preserve the functional integrity of the liver

PHIORITZIN DIABETES

So-called 'phlorhizin diabetes" was discovered and first described by von Mering in 1879 (12). It results from the administration to experimental animals of the

Table 10*

Comparison of the Effects of Loutropic Factors
(Micherry and Patterson [45])

Regimen Used for Production of Patty Livers	Chal ne	Lipocase	Inca tol
Departrestized dogs	++*	++*	-7
Rath Bigh fat diet Talumine All B vitamine All B vitamine Electrical Talumine Thampe Thampe Tham ne and riboffavan Thamme choffavan pymfonne and pastothenic scid Above four vitamin and b othe B vitamine and obselveroid	###	++111 010	114 0177

[&]quot;Symbols ++ strang" potropic action + made are ligatropic action a so ispotragic action - initiate " verified in two of more laboratories

glucoside phlorhizin (or phlorhidzin), which has the structure indicated in Figure 27. The drug is generally administered subcutaneously, as a fine suspension in oil, and the issual dosage is about 1 gm of phlorhizin per day for a zo-Lg. dog. (xg)

Fig 27 -Phlorhuin

In order to obtain a rapid initial effect the first dose is sometimes administered in a 2 5 per cent sodium bicarbonate solution (14)

The syndrome of phlorhizin diabetes (15) and its progression to the death of the animal resembles that of pancreatic diabetes in practically every particular ex

cept that the blood sugar level is abnormally low (hypoglycemia), as opposed to the hyperglycemia of the deparcreatized animal As has been previously indicated (D 77), the drug produces its effect by preventing the reabsorption of sugar by the tubules of the Lidney This is accomplished by the inhibition of the phosphory lation of the glucose to hexosephosphate (16) All tissues are subject to the same action of phlorhizin But muscle tissue destroys phlorhizin very quickly, so that ef fective concentrations of the drug in muscle are not attained by the procedure em ployed in producing phlorhizin diabetes in the living animal (17, 18) However, under an vitro conditions the action of phlorhizm on isolated muscle tissue can be readily demonstrated (19) As used in prio, the kidney shows the greatest effects of phlorhizm because it has a limited ability to destroy the drug (17) and also because the excretory function of the kidney leads to the accumulation of phlorhisin in larger concentration than elsewhere in the body (15) Hence, phlorhizin dia betes may be regarded as being primarily a disturbance in the kidney. This was shown at an early date by Minkowski, who demonstrated that the removal of the kidneys from phlorhizinized dogs abolished all signs and symptoms of diahetes during the time of survival of the animals in the absence of renal excretory func tion (II)

A comparison of pancreatic and of phlorhizin diabetes indicates that the poly uria, polydipsia, dehydration and demineralization, loss of weight, weakness and polyphagia, and ketosis and come are dependent, in both, on the loss of significant quantities of carhohydrate from the body by way of the urine. In pancreatic diabetes, this results from a disturbance in the regulation of the blood sugar, leading to hyperglycemia, which, in turn, exceeds the capacity of the phosphorylative mechanism of the kidney for the reabsorption of sugar. In phlorhizin diabetes, the same train of events is initiated by a lowering of the phosphorylative capacity of the kidney, allowing a significant excretion of sugar at normal and hypoglycemic blood sugar levels.

THE NON UTILIZATION THEORY OF DIABETES

During the ten years that followed the discovery of pancreatic diabetes by von Mering and Minkowski in Strassburg, the same laboratory established the classical criteria of the metaholic disturbance in experimental diabetes (20). These criteria comprise (1) the quantitative excretion of administered carbohydrate in the unne of the experimental animal, (2) the unnary destrose to nitrogen ratio (D. N.), (3) the excretion in the unne of acetoacetic acid, β hydroxybutyric acid, and acetone (ketosis), and (4) the characteristic respiratory quotient (R.Q.)

The quantitative excretion of administered sugar by the diabetic animal suggested that the cause of the metabolic difficulty was an mainlify to utilize carbohydrate (the non utilization theory). Furthermore, when Minkowski collected urms specimens from his deparcreatized dogs (while lasting or when fed lean

meat) and analyzed them for amounts of dextrose and netrogen, respectively, the total amount of sugar in each 24 hour specimen seemed to bear a definite relationto an amount of introgen in the same spectmen (6, 11). This D. N ratio aver says to the amount of metogen at the same spectmen $\{0,11\}$ AMS J it ratio averaged about 2.8. I for his animals (see Table 11), from which he concluded as follows:

ORIGINAL DATA OF MINEORSEI (41) OV SLOAR AND NITEOGEN EXCRETION OF DEPARCREATIZED DOOS

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eared nitrogen		i 12 4	5 95 3 76	
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lows (a) Since nitrogen is a breakdown product of protein, all the sugar which ap nows (a) since mitrogen as a measurowin product on protein, an inc nogar which appeared in the urine was being made at the expense of protein (b) From the apparpeared in the utime was being made at the expense of protein (v) a run the apparent containey of the D N ratio, none of the sogar made from protein was being out constancy or the way fails, hone or the popul made from process withined by the diabetic animal, i.e., all of it was quantitatively exercted

The appearance of the ketone bodies in the diabetic animal was the third basis for the non utilization theory of diabetes. It was known that acctoactic acid and δ bydroxybutyric acid resulted chiefly from the breakdown of fat Since these substances did not ordinarily appear during fasting in the normal organism (when fat was the chief metabolite), it was assumed that the ketone bodies were abnormal waste products resulting from the incomplete exidation of fats in diabetes. From this arose the conception that a certain amount of carbohydrafe had to be oxidation or derivating from the incompletely ("fats burn in the fire of carbohy drates") (21, 22, 23). Thus the ketoes of diabetes was apparently another evidence of the lack of ability to utilize carbohydrafet.

Studies of the respiratory exchange of the normal and diabetic organism apparently supported the foregoing conclusions. If the net result of complete oxidation in the body is compared to the burning of a substance in a bomb calonimeter, it is apparent that the amount of oxygen consumed and the amount of CO, given of in the process will depend upon the chemical nature of the substance that is boing oxidized. Thus it may be calculated that, when a carbohydrate is conduzed, I mol of CO, will result for every mol of oxygen used, according to the reaction

The R Q is the relation expressed in volumes, between the oxygen consumed and the CO₂ given off (CO₂/O₃). Hence the R Q for the oxidation of carbohydrate is to In the same way, it may be calculated that the R Q for fat is about o 7, for protein, about 0 8. The latter figure involves a number of assumptions, since protein is not entirely oxidized in the living organism (24, 25).

It was found that the R Q of a normal animal under fasting conditions was in the neighborhood of o 7 lbis was taken to indicate that fait was the chief fuel being used at that time. After a carbobydrate meal the R Q of the normal animal rose toward to (Fig. 28). This was interpreted to mean that the animal was now oxidizing the ingested carbobydrate. The diabetic organism differed from the normal in that, while its fasting R Q was also about o 7, the quotient did not hewhen carbobydrate was administered (Fig. 28). This seemed to confirm the conclusion that the diabetic organism cannot use carbobydrate but derives its energy chiefly from fat (4a, 26, 7).

A CRUCIAL EXPERIMENT OPPOSING THE NON UTILIZATION THEORY OF DIABETES

On the basis of the four lines of evidence which have been outlined, the non with zation theory of diabetes was more or less generally accepted for many years. This was made possible by ignoring certain inconsistencies in the evidence and by neglecting other evidence to the contrary. As early as 1897, Kausch (28) reported the results of removal of the liver from depancreatized geese and ducks, as com

pared to the results of the same procedure in normal birds. He found that, in the absence of the organ which supplies the blood sugar, the latter disappeared from the blood just as quickly in the diabetic birds as in the normal ones. There were a number of subsequent attempts to confirm this finding in mammals. Most of them showed similar results (29 30, 31), but technical difficulties as regards complete removal of the liver and the consequent irregulantly of the data rendered these findings inconclusive.

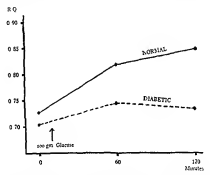


Fig. 28—Rise of R Q following sugar administration to normal and departmentized dogs. (From the data of Barker et~al. [26].)

However, following the development of Mann's technic for total removal of the liver in dogs Mann and Magath (37) reported unequivocal evidence that the completely depanceatized dog suffers just as rapid a fall in the blood sugar after hepatectomy as does the normal dog (Fig. 29). Whether originally normal or dia bette, the liverless animal dies in hypogly cours convulsions within a few hours. In either case it can be kept alive only by continuous administration of sugar or the giving of larger amounts of sugar at about 2 hour intervals. Unless one makes the rather absurd assumption that the removal of the liver suddenly restores the ability of the peripheral tissues to utilize carbohydrate, one must conclude that the diabetic animal does not lack that ability. Under these circumstances it be-

comes important to re examine the classical criteria of diabetes for their true mean ing and to consider all other evidence which may help to explain the diabetic syn drome without invoking the non utilization theore.

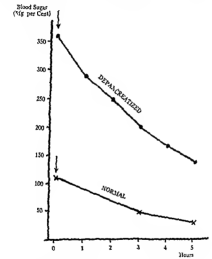
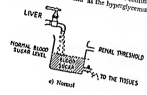


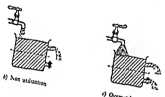
Fig. 2g — Development of hypoglycemta folior mg hepatectomy in depanceatized as well as in nor mai dogs. (Mann and Magath 134)

THE OVERPRODUCTION THEORY OF DIABETES

The alternative to the non initization theory of diabetes is the overproduction theory of diabetes. These two possible explanations for the diabetic syndrome are compared in Figure 30 in terms of a simple mechanical analogy. Diagram A indicates the state of affairs in a normal animal in which the liver, as represented by

the tap, 13 pouring just as much sugar into the blood as the tissues (represented by the lower, outflow tithe) are drawing off for utilization. The net result of the dy the tower, outflow time) are grawing on for utilization. The net result of the dynamic balance between inflow and outflow is the normal blood sugar level. Dis gram B represents the non utilization theory adopted by Minkowski Here the gram o represents the non numerous theory adopted by attractions and the tissues has crased while the liver continues to pour sugar outdown up sugar into the tissues has created white the first continuous of processing approaches the





c) Overproduction Fig. 30 - Mechan cal analogy allustrating the alternative theories of diabetes

renal threshold glycosuna begins Diagram C represents the other possible ex tenal intesting glycosum begins traggator compressed to other prostore ex-planation, first proposed by von Noorden (33) and later advocated by a vigorous premature must proposed by you arounce 1337 and later advocated by a vigorous minority (34–35–36), namely, the overproduction theory. Here, there is no dimi minority (34-35-30), namely, the overproduction theory. Here, there is no dimination of the utilization of blood sugar by the tassies. But the supply of sugar to nation of the utilization or oboos sugar by the cossues over the supply or sugar to the blood from the liver has become excessive to the point where continued normal the mood from the to et has become excessive to the point where continued normal stillization can no longer keep pace with it. Hypered crims and glycosuma follow Figure 30 makes it obvious that closing the tap (hepatectomy) would produce A feart 30 makes it outloop that cooming the cash (hypogly central) in diagrams A

and C but not in diagram B Thus, while both theories can account for cardinal features of the diabetic syndrome, the non utilization theory is directly opposed by the hypoglycemic effect of hepatectomy in the diabetic animal (p. 97). There is no conflict in this regard if one adopts the overproduction theory. The re examination of the classical criteria of diabetes which is the subject matter of the subsequent three chapters should, therefore, be followed with reference to both the possibilities indicated in Figure 30.

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PART III CRITICAL SURVEY OF THE CLASSICAL CRITERIA OF DIABETES



QUANTITATIVE EXCRETION OF ADMINISTERED SUGAR AND THE DEATROSE NITROGEN RATIO

HE fact that the administration of destrose to his diabetic animals to HE fact that the administration of dextrose to his diabetic animals resulted in the exerction of a roughly equivalent amount of sugar in the suited in the excretion of a roughly equivalent amount of sugar in the control Eight 30 (p 99) will show at a glance that his conclusion was not a logical access. Figure 30 (p. 99) will show at a glance that his conclusion was not a logical necessity. It may be seen that viewed from the standpoint of either theory, the influx sily It may be seen that viewed from the standpoint or either theory, the innur of angar into the blood would be expected to result in an entra of an extra amount of sugar into the blood would be expected to result in an extra amount of sugar in addition to that which is already over dowing through the kidneys

DEXTROSE NUTROGEN RATIO IN THE DEPANCREATIZED ANTHAL It is important to consider to detail the supposed constancy of the D N ratio It is important to consider to detail the supposed constancy of the D. A. Action.

If it mere truly constant, it would constitute strong support for the non-ruthleation strong support for the non-ruthleation. It it were truly constant it would constitute strong support for the non utilization for a rate of sugar utilization (other theory for it nould be difficult to conceive of a rate of sugar utilization (other than zero utilization) so univarying in different diabetic animals and under different diabetic animals and under different different different dispersions. than ero utilization) so unvarying in different diabetic animals and under diabetic animals animals and under diabetic animals and under diabetic animals animals animals animals and under diabetic animals and under diabet ent conditions as to make the ratio possible. Minkowski s summarized data are re-Produced in Table 11 (P 95) On 31 experimental days in 9 departmental days in 9 departmenta led on meat be obtained D N ratios which valved from 3 to 1 to 2 for 1 with an average of 2 8 1 (1) The experimental days which he used to establish the average ayrage of 2 8 f (1) The experimental days knich he used to establish the average and the control of the control of all the experimental days on any ratio are admittedly selected since a record of all the experimental days on any interest and a selection of the experimental days angle animal round show D N ratios much augner than 2 & 1 to percury with and the fallow which fell progressively below this figure as the exitus of the animal was also ratios which fell progressin ely below this figure as the exitus of the anuscal was approached. The high initial D N ratios were discarded on the basis that they approached the nigh initial D. A ratios were discarded on the basis that they represented the pouring out of preformed circovers stores. The low D. N. ratios represented the pouring out of preformed given stores. The low D N ratios of the experiments were disregarded because of the poor condition toward the end of the experiments were disregarded because of the poor condition of the animals at that time. The reasonableness of these objections to the results of the animals at that time. The reasonableness of these objections to the results that the days of each experiment cannot be desired. But a closer exof the first and last few days of each experiment cannot be denied that a closer ex
amination of Vinkowski s data mater it apparent that the experimental days were

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the state of the experimental days were

th amnation of Minkowski s data makes it apparent that the experimental days were selected in a much more arbitrary manner than we have been led to believe by selected in a much more arbitrary manner than we have been led to believe who have trustingly accepted his average as a physiological constant

hose who have trustingly accepted his average as a physiological constant.

The analysis in Table 12 shows that the selected data in any five experimental and the selected data in any five experimental and the selected data in any five experimental selection. The analysis in 12016 12 shows that the selected data in any given experimental began as early, as the second day of disbetts or as late as the twelfth day animal began as early as the second day of diabetes or as late as the twelfth day of diabetes or as late as the twelfth day

Moreover, the days reported in some experiments are not consecutive, some days being omitted, for no stated reason. It must be apparent that any desired average D N ratio might have been obtained by such arbitrary selection of experimental days, picked from experiments in which the D N ratios fell progressively from high to low values.

This criticism is supported by other results of Minkowski—reported in the same paper but not included in the figures from which be obtained his D N ratio of nomparing the initial D N ratios obtained from well nourished and pooly nour ished animals he recorded ratios in the latter animals of 2 04, 2 43, 1 62, and 2 24 on the third, fourth, and fifth days of diabetes. It is difficult to understand why Minkowski tid not attempt to correlate these low results with the data from

TABLE 12
THE DAYS DURING THE DIABETIC LIFE OF HIS DOOS WHICH MINKOWSKI (r)
USED TO COMPUTE HIS AVERAGE D N RATIO

Dog No					Day	s afte	r Panc	reblec	tomy					
I II IV VI VII VIII VIII IX	2	3	4	3	6	7	8	9 9	10 10	11	13	13	14	15

which he computed his average ratio. The poor nutrition of these animals might perhaps have accounted for the failure to obtain high initial D. N ratios. But the values uniformly below a 8.1 obtained on days in which the approaching demais of the animal was not a factor and on days which coincided, in point of time, with some of the experimental days which were used to obtain his average, serve to con firm the arbitrary nature of the average D. N ratio at which he arrived. This indication of the inherent defects in Minkowski, s work is not intended to cast aspersions on his integrity as a physiologist. It must be remembered that Minkowski, working before the days of insulin, had to deal with actuely diabetic dogs suffering from the effects of a recent anesthetic and operation.

Pflueger (2), Embden (3), and others subsequently reported that they had failed to obtain fixed D N ratios at the Minkowski level Their work was criticized on the assumed ground of the poor condition of their animals or of incomplete pancreatectomy. Such criticism, however, cannot be leveled at the work of Mac leod and Markowitz (4), who used depancreatized dogs that were maintained in excellent condition by the use of insulin After the withdrawal of food and insulin from such animals (which by subsequent post mortem examination were shown to

be completely deparamentatized) they obtained D N ratios far below 28 1, after be completely departeratized) they obtained D N ratios far below 28 1, after that departed Charloss and co-workers (3) re the first few days of the experiment had elapsed Unaskott and co-workers (3) reto-complete and found (as noted by Minkowski) that the D N ratio was ported similar results and found (as noted by Minkowsky) that the D. N. ratio was a decidedly in the generally higher in fat than in lean does and also that it varied decidedly in the same animal according to its autintsonal condition at the time of the experiment and animal according to its autotional condition at the time of the experiment at the time of the Experiment to the D V ratio in ad In 1930 Rapport (6) reviewed the extensive interasting on the D V ratio in addition to the above and was not able to reconcile the large variations which had to, ation to the above and was not able to recordie the large variations which had been reported. In the same year Sockin (f) published a comprehensive remirch had been reported by the same year Sockin (f) published a comprehensive remirch had been reported by the same year Sockin (f) published a comprehensive remirch had been remired by the same year Sockin (f) published a comprehensive remirch had been reported by the same year Sockin (f) published a comprehensive remirch had been reported by the same year Sockin (f) published a comprehensive remirch had been reported by the same year Sockin (f) published a comprehensive remirch had been reported by the same year Sockin (f) published a comprehensive remirch had been reported by the same year Sockin (f) published a comprehensive remirch had been reported by the same year Sockin (f) published a comprehensive remirch had been reported by the same year Sockin (f) published a comprehensive remirch had been remirched by the same year Sockin (f) published a comprehensive remirch had been remirched by the same year Sockin (f) published a comprehensive remirched by the same year Sockin (f) published a comprehensive remirched by the same year Sockin (f) published a comprehensive remirched by the same year Sockin (f) published a comprehensive remirched by the same year Sockin (f) published a comprehensive remirched by the same year Sockin (f) published a comprehensive remirched by the same year Sockin (f) published a comprehensive remirched by the same year Sockin (f) published a comprehensive remirched by the same year Sockin (f) published a comprehensive remirched by the same year Sockin (f) published a comprehensive remirched by the same year Sockin (f) published a comprehensive remirched by the same year Sockin (f) published a comprehensive remirched by the same year Sockin (f) published a comprehensive remirched by the same year sockin (f) published by the been reported in the same year Nosten (1) punished a comprehensive reinvestiga ton of the D N ratio in depondreatured dogs using the advanced feedingue made ton of the D N ratio in deparcratized dogs using the advanced technique made to the discovery of maining. This work was done on deparcratized dogs. Possible by the discovery of insuling This work was done on departmentation dogs the use of insuling and present

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ed well besied and non infected wounds The animals were maintained on a low ed well healed and non injected wounds the animals were maintained on a low Caloric Protein diet and the absence of solet 1880e was verbed by post morton exannation in contrast to Almkowski's summer they survived the withdrawal or so long as 5 weeks during which time they usually remained bright and insulin for at long as 5 weeks during which time they usually remained briefst and the D. N. ratio comprised 138 active although loung aright. The observations on the U. N. fatto comprised 133 selected days for 50 dogs in contrast to Minkowski 5 31 selected days for 9 dogs. Unsolected days for to dogs in contrast to Minkowski 5 31 selected days for 9 dogs of the D N Values obtained is shown in Table 53 It may be The distribution of the D N Values obtained in shown in Table 13. It may be a contained and though some D N values and at 10 Mink one 14. It may be a contained the contai seen that although some U N tatios similar to almhossast's arre-collained there is nothing to indicate that such values have any particular againscance in general to nothing to indicate that such values have any particular segmentace in general segments. At a state the beginning of each experiment and to show are D. V fatto tended to be high at the beginning of each experiment and to show a progressive fall as the animals lost weight and their stores of adaptive times were a progressive fall as the animals lost a cight and their stores of adipose tissue were constant to a cight and their stores of adipose tissue were constant as the animals manufactured by processing the cight and the processing the cight and their stores of adipose tissue were constant and the cight and their stores of adipose tissue were constant and the cight and their stores of adipose tissue were constant and the cight and their stores of adipose tissue were constant. depleted. This serves to explain the observed D V ratios reported by previous maintained D N values far below 20 1 for WORLETS The fact that some sammals maintained D N values for below 2 0 1 for home and an analysis of the appellation of premoral which some writers of all colons and analysis of the appellation of premoral which some writers as long as 16 days precludes the appellation of Primortal which some with the Minkowski level (8) Ave used to at old consideration of all raties before the atmixorals let et (8) It is clear that it Minkowski s interpretation of his ratio is accepted the progreatively lower ratios obtained later in the experiments agonly the villecation of amounts of the sugar arising from protein If on the other hand the

low ratios obtained later in the experiments represent the true extent of gluconeogenesis from protein, the higher Minkowski values must mean that sugar is being formed from fatty acid as well as from protein. In either case, there remains no basis for concluding that sugar is derived solely from protein or that none of the sugar so formed is utilized by the diabetic organism. It is permissible to conclude that sugar is derived partly from protein, but it is impossible to say to what extent this occurs.

DEXTROSE NITROGEN RATIO IN THE PHLORHIZINIZED ANIMAL

Conclusions similar to those arrived at with respect to pancreatic diabetes may be drawn in regard to the significance of the D N ratio of 3 65 1 obtained by some investigators in so called "photohizm diabetes" There is an added difficulty in in terpreting this type of work in that there is no standard for judging the experimental preparation, comparable to the histological demonstration of complete pancreatectomy in operated animals I it is obviously fallacious to account for different D N ratios obtained with phlorhizm in different animals and by different workers (13) by saying that some of the animals were not completely phlorhizm itzed hecause they did not yield D N ratios of 3 65 1 An added completation is the fact that the phlorhizm, as used, is not a pure chemical substance of known composition In his last gublication on the subject, Graham Lusk (3) (who together with his coworkers had made the most extensive use of phlorhizm diabetes in their studies) confessed that with the phlorhizm he was then able to obtain he could not reproduce the D N ratio of 3 65 1 which he had formerly insisted was the necessary criterion for complete phlorhizmination.

Even those workers who used preparations of phlorhizin with which they were able to obtain some D N ratios approximating 3.65; I were not able to maintain such ratios at will in a given animal A so in the depancreatized organism, the D N ratio resulting from continued phlorhizin administration starts at a high value and declines progressively. The selection of days upon which the ratio is to be considered a valid one is a purely arbitrary matter. Table 14 and Figure 31 show the day by day excretion of sugar and introgen in the urine and the D N ratio in three dogs receiving the customary phlorhizin treatment. It may be seen that there is no evidence for a constant D N ratio at any level.

If, for the moment, one were to discount the foregoing considerations, one would still have to explain the difference between the phlorhizm D N ratio of 3 6 3 1 and the Minkowski ratio of 2 8 1. There is no factual basis for concluding that phlor biant alters the biochemical processes in such a manner as to allow a larger proportion of the protein molecule to be converted into sugar And, if a constant proportion of the protein molecule is convertible, one or both of the following conclusions is unstified either the depancreatized animal always utilizes a significant fraction

of the sugar derived from protein or the phlorhizmized animal must be forming sugar from fat as well as from protein Finally, one must take into account the fact that even the classical criteria are

Finally, one must take into account the fact that even the classical criteria are self contradictory as regards the ability of phlorhizmized animals to utilize carhohydrate

 a) Insulin has been obtained from the pancreas of dogs after prolonged and maximal phlorhizin treatment (ro)

TABLE 14

LACE OF CONSTANT D N RATIO IN FASTED PHLORHIZINIZED DOGS

Dos No	LENGTH OP EXPT	Urner En (Gr 72		DN	Union Revones	BLOOD
	(DAYS)	Destrose	N troges		LETONES	(Mo res Cert)
ī	2 3 4 5 6 7 8	16 05 11 16 9 36 6 22 2 26 3 80 4 86 2 85	5 95 4 43 3 70 2 89 1 73 2 83 2 84 2 72	2 69 2 52 2 53 2 15 1 31 1 35 1 43 1 05	++ +++ +++ ++ ++ ++ ++	14 21 18 26 14 12 31
,	1 2 3 4 5 6 7 8 9	6 46 \$ 55 3 69 4 69 4 89 2 51 2 51 1 87 3 15	7 31 3 11 1 94 2 36 2 36 1 86 1 56 1 40 2 09	2 79 1 79 1 99 1 73 2 07 1 35 1 65 1 26 1 50	0000++	38 33 33 20 11 17 31
3	1 2 3 4 5 6 7 8 9 IO	24 61 15 86 13 18 11 00 0 89 8 11 9 55 8 13 7 92 5 49 5 14	7 84 6 58 5 86 6 41 5 23 4 77 4 52 4 40 4 61 3 82 4 26	3 14 2 25 1 71 1 89 1 70 2 11 1 85 1 73 1 44 1 23	++ ++ ++ ++ ++ 0 ++ + 0 0	30 29 24 25 22 18 21 21 25 21 20 24 25 26 27 28 29 29 20 20 20 20 20 20 20 20 20 20 20 20 20

b) After nephrectomy the phlorhuzanzed dog is quite normal as regards its blood sugar level its R Q and the rise in the R Q following glucose administration (11, 12)

c) The ingestion of sugar by the intact phlorhizmized animal results in the retention of glucose which has an anti-togenic and protein sparing action, and causes a rise in the R Q comparable to that which occurs in the normal dog [11, 13, 14, 15 16].

CHAPTER X

KETOSIS

F THE three substances usually grouped under the term ' ketone bodies namely, acetoacetic acid, \$\beta\$ bydroxybutyric acid, and acetone the second is not a Letone, and the third represents merely a breakdown product of its more physiologically significant precursors. It is now generally agreed that, under conditions leading to ketosis, acetoacetic acid is the first ketone body to be formed (1) It is known that various tissues of the mammalian organ ism are able to reduce acetoacetic acid to β bydroxybutryic acid and also to effect the reverse reaction The direction of this reversible reaction depends on the con centration of substrates present and on the oxygen tension, and there is evidence that an equilibrium between these two substances is established rapidly (2, 3, 4) Hence it is a matter of practical importance, in balance or recovery experiments, to estimate the amounts of both of these substances present in the tissues when at tempting to account for the fate of a given amount of either. Acctone is readily formed in solutions containing acetoacetic acid, and it is generally assumed that whenever it is found in biologic fluids it is merely a spontaneous decomposition product which indicates that an equivalent amount of one of the other ketone bodies was formerly present

SITE OF ORIGIN OF THE KETCHE BODIES

Practically all investigators have agreed as to the chief source of the ketone

bodies A similar conclusion regarding Letogenesis by these organs in situ was reached by Himwich, Goldfarb and Weller (9 10), who compared the ketone levels of the inflowing arterial blood and of the outflowing venous blood of the variation of the control blood for the control b

the reverse

sional output of small amounts of ketone bodies from the skeletal muscles and the intestinal tract. In agreement with this, Jowett and Quastel (11) found that slices of kidney, spleen, testis, and brain in vitro could produce small amounts of the ketone bodies from butyric acid but that liver slices under similar conditions produced from ten to forty times as much.

It should be noted that the evidence quoted above does not prove that organs other than the liver are incapable of forming considerable amounts of the ketone bodies For it is obvious that when a tissue is capable of utilizing a substance, the amount of the latter which may escape from that tissue into the blood for surrounding medium in tirbo is merely the difference between the amount formed and the amount utilized in situ. That this is not a theoretical consideration only was shown by Weinhouse (63) for kidney tissue, using the heavy carbon tracer technic Under these circumstances: the role of the liver as the chief site of origin of ketone bodies depends upon the fact that it can form these substances at a much greater rate than it can utilize them.

Whether or not the extrahepatic tissues can be shown to put out some ketone bodies under special experimental conditions, it is clear that in the living infact animal the liver is practically the sole source for these substances. Thus it has been demonstrated that dogs in which the functional capacity of the liver is limited by an Eck fistula do not exhibit increased ketosis after phlorhizm administration (12). The reduction of bepatic function by hepatotous agents also decreases

the diabetic animals was due to rapid ketogenesis in the liver. Finally, Mirsky (15) has recently shown that the ketogenic effects of certain pituitary, extracts, which are regularly obtained in normal animals, cannot be demonstrated in the absence of the liver.

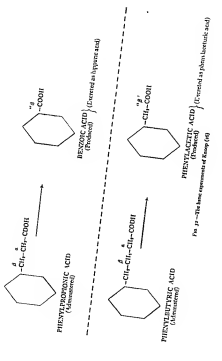
SOURCE MATERIALS FOR PRODUCTION OF KETONE BODIES

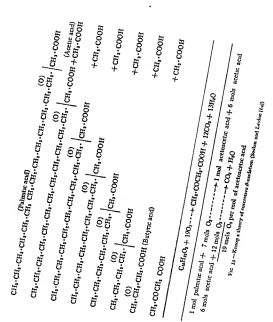
The early work of Embden and co-workers (16 17) indicated the formation of extra ketones by perfused livers when fatty acids certain amino acids or pyrivic acid were added to the perfusing fluid. These three different source materials for the Letone bodies have since been confirmed by a number of investigators in a variety of ways (18 19 20 21 22 23) However, Embden and associates reported that the amount of ketone hodies arising from fat greatly exceeded that from the other sources Subsequent work has emphasized the fact that, when ketosis occurs in the living organism it may be regarded for practical purposes as an index of the catabolism of fat Thus the perfused fatty liver produces much greater amounts of the ketone bodies than the liver that is poor in fat (23) The livers of depancreatized or phlorhizmized animals which are characteristically rich in fat, are known to produce excessive amounts of Letone bodies (22) In the intact nor mai animal the feeding of fat or the excessive use of depot fat, induced by starva tion, results in ketosis. More recently. Stadie Zapp, and Lukens (24 25) have demonstrated that the production of ketones by liver shees in trito is accompanied by the disappearance of amounts of fatty acid sufficient to account for more than I mol of Letone per molecule of fatty acid

For many years the general conception of the mechanism by which ketones are formed from fatty acids seemed to be quite settled, but it has recently undergone at least two metamorphoses The theory of successive \$\beta\$ oxidation originated from the work of Knoop (26) It was based on the feeding of various phenyl substituted fatty acids to test animals and the identification of the excretion products in the urine The administration of either benzoic, phenylpropionic, or phenylvaleric acid resulted in the appearance of hippuric acid. After the administration of phenyl acetic and phenylbutyric acids, phenylaceturic acid appeared in the urine (Fig 32) These results could be reasonably explained only by assuming that the fatty acids were degraded by the splitting off of two carbon atoms at a time, by orida tion at the carbon atom which occupied the \$ position in relation to the carbonyl group It was assumed that the acetic acid molecules so formed were rapidly metab olized, while the phenyl group was left attached to one or two carbon atoms, de pending on the original number of carbon atoms in the fatty acid molecule This assumption was confirmed in pito by Dakin and was extended to the in titro oxida tion of various fatty acids by hydrogen peroxide at body temperature (27, 28 29) Snapper, Gruenbaum, and Neuberg (7) duplicated Knoop's results on the per fused kidney

With this groundwork laid Embden and co workers (5, 6) perfused various fatty acids through isolated livers and reported that ketones were formed from fatty acids with an even number of carbon atoms in the molecule but not from the odd numbered fatty acids. This confirmed the natural occurrence of β oxidation and also seemed to indicate that the last four carbon atoms in the chain under went oxidation at the \$ position but were not split It was, therefore, assumed that each molecule of an even numbered fatty acid, regardless of chain length, resulted in the production of one molecule of ketone and that odd numbered fatty acid could not give rise to ketone bodies. On this basis, also, the amount of oxygen re quired for the degradation of a given fatty acid and the production of one molecule of ketone could be calculated (Fig. 33)

Although this conception gained wide popularity (especially among clinicians concerned with clinical states characterized by ketosis) and although it persists in many textbooks up to the present day, serious objections from the experimental standpoint arose before many years had passed Thus, Hurtley (30) sought for the butyric and acetic acids that would be expected to be present in the liver during active ketogenesis and failed to find them Clutterbuck and Raper (31), Smedley-MacLean and associates (32, 33), Witzeman (34), and Verkade and van der Lee (35), who repeated and extended the in vitro work of Dakin, found that, while β-orndation did occur, oxygen could also become attached at the a and the γ posi tion A more serious objection, from the point of view of the whole animal, was the observation by Deuel and associates (36, 37) that more ketone bodies arose in an





KETOSIS

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animal fed octanoic acid (Ca) than in an animal fed an equimolar amount of butyric acid (C4) Shortly afterward, Jowett and Quastel (1, 11), and later Leloir and Muñoz (21), observed that the amounts of ketone hodies formed by liver slices in vitro could not be accounted for on the assumption that only the last four carbon atoms of each fatty acid molecule gave rise to a ketone body. A similar discrepancy was reported for perfused livers by Blixenkrone Møller (38, 30) and for liver slices in titro by Stadie and co-workers (40) when the oxygen consumption during experiments was compared with that which would have been expected on the basis that all but the last four carbon atoms of each fatty acid was being dis posed of by the oxidation of the acetic acid formed. The observed oxygen consumptions were far smaller than would allow for this mode of fatty acid breakdown Finally, the improved technies for ketone estimation, which have made possible the determination of relatively small amounts in blood and tissue, bave led to the recent finding that the odd numbered fatty acids also give rise to smaller but significant amounts of the ketone bodies, as compared with the even numbered fatty acids This has been reported by Jowett and Quastel (x, xx), Edson (Ax). and Leloir and Muñoz (21) for isolated tissue (liver) and by MacKay and associates (42) for the intact animal

It is obvious that the hypothesis of successive \$\tilde{\text{D}}\tilde{\text{Sundation}}\$ in the aforementioned form is no longer tenable. Indeed, as long ago as 1916, Hurtley (20) proposed the theory of multiple alternate exidation to account for his failure to find butyric and acetic acids in ketone producing hiver. He expressed the opinion that the intact fatty acid chain was first outliered at each alternate carbon atom and then spht into blocks of four carbon atoms each—a process which would not necessitate even the transiert presence of either of the substances for which be tested. According to this hypothesis, the number of ketone molecules arising from a fatty acid would be the whole portion of the quotient when the number of carbon atoms in the fatty acid molecule is divided by 4. This hypothesis was adopted by Deuch, Quastel, Leloir, Blixenkrone Mfeller, and Stadie, since it accounted for the greater than 1 x ratio of ketogenesis from the higher fatty acids, the lower oxygen consumption than that expected from the x x ratio, and the formation of ketone bodies from odd numbered fatty acids (Fig. 34). Until recently the multiple alternate oxidation theory was adequate to explain.

the available data. However, it implied a phenomenon rather difficult to explain on biochemical grounds. The simultaneous candation of every alternatic carbon atom offered no difficulty. But how could one explain the selective splitting of the molecule at every sector date for group instead of at every keto group? This difficulty is avoided by a newer conception, which also accounts for other recent evidence not compatible with the theory of multiple alternate oxidation. In a systematic interesting of the ketogenic properties of fairly acids consisting of from one to eleven carbon atoms. Jowett and Quastel(1,11) noted, among other things, ketone pro

CH ₁ ·CH ₁ ·CH ₁ ·CH ₁ ·CH ₂ ·CH ₂ ·CH ₂ ·CH ₂ ·CH ₂ ·CH ₃ ·CH ₃ ·CH ₃ ·CH ₃ ·COOH + CH ₃ ·COOH	(Acetic acid) H +CH ₄ ·COOH
CH,	+сн,-соон
(O) (CH ₁ ·CH ₁ ·CH ₁ ·CH ₁ ·CH ₂ ·CH ₂ ·CH ₂ ·CH ₂ ·CH ₃ ·C	+сн, соон
(O) CH-CH-CH-CH-CH-CH- CH-CH-CH-CH-CH-CH-CH-CH-CH-CH-CH-CH-CH-C	+сн,соон
CH-CH-CH-CH-CH-CH-COOH	+СН•СООН
(O) CH1-CH1-COOH (Butyne acid)	+сн, соон
CHCOCH-COOH	
C _a H ₃ O ₃ + 19O ₃ → CH ₃ -COCH ₄ -COOH + 12CO ₄ + 13H ₄ O	
1 mol paintic acid + 7 mols O+1 mol acetoacetic acid + 6 mols acetic acid 6 mols acetic acid + 12 mols O ₂ +CO ₂ +H ₂ O 19 mols O ₂ per mol of acetoacetic acid	ols acetic acid

(Palmitic acid) (O) (CH₁·CH₂·CH₂·CH₃·CH

animal fed octanoic acid (C) than in an animal fed an equimolar amount of buannua ieu ocianore acin (Ca) man in an annua ieu an equinnosar amount oi oci-tyric acid (Ca). Shortly afterward, Jorett and Quastel (1, 11), and later Leloir and Greecia (A) anorthy atterward, Junetic and Quaster (1, 11), and later Lemmand Muñoz (21), observed that the amounts of ketone bodies formed by liver slices Author (21), observed that the amounts of actione occurs formed by fiver successions that only the last four carhe time could not on accounted for on toe assumption that only the last rour carbon atoms of each fatty acid molecule gave rise to a ketone body. A similar disoog atoms of each satty acm morecuse gave use to a setone oody. A similar dis-crepancy was reported for perfused livers by Blatenkrone Møller (38, 39) and for crepancy was reported for pertused avers by differentiations support (10, 39) and to the silves in title by Stadie and co-workers (40) when the oxygen consumption aret suces in time by stable and co-workers (40) when the oxygen consumption during experiments was compared with that which would have been expected on curing experiments was compared what that which would have been experied on the basis that all but the last four carbon atoms of each fatty acid was being dis the outs) that an out the last tout cations around ve each tarty actu was being this posed of by the oxidation of the acetic acid formed. The observed oxygen con poses of by the explantion of the access acres acres for this mode of fatty and break sumptions were tar smaller man would allow for this mode or fatty acid areas down. Finally, the improved technics for ketone estimation, which have made toom rmany, the improved technics for ketone estimation, which have made possible the determination of relatively small amounts in blood and tissue, have possume the determination of relatively small amounts in blood and tissue, bave ided to the recent finding that the odd numbered fatty acids also give use to smaller texto use recent undust that the out numbered latty, acids also gave rise to smaller but significant amounts of the ketone bodies, as compared with the even numbered out significant amounts of the xerone poures, as compared with the even numbered fatty acids. This has been reported by Jovett and Quastel (1, 11), Edson (41), native actus 1 ms uss peen reported by Jowett and Quaster (1, 11), Edson (41), and Musics (21) for isolated tissue (liver) and by MacKay and associ ates (42) for the intact animal

to (42) for the intact annual II is obvious that the bypothesis of successive β -ordation in the aforementioned at is our rows that the hypothesis of successive postulation in the autorimentioned form is no longer tenable. Indeed, as long ago as 1916, Hurtley (30) proposed the torm to no tonger tensore tinuecu, as long ago as 1910, fluttey (30) proposed the sheety of multiple alternate exidation to account for his failure to find butyric and acoty or multiple difference outdation to account for the similar to find outsylve and coin acids in Actione producing livers. He expressed the opinion that the intact cent actus in actume producing avers are expressed the opinion that the infact ity acid chain was first oxidized at each alternate carbon atom and then split the control of the control of the carbon atom and then split tty acid enain was rist oxidized at each anternate caroon atom and then sput to blocks of four carbon atoms each—a process which would not necessitate even to mocas of four caroon atoms eaco—a process which would not necessitate even a transient presence of either of the substances for which he tested. According s hypothesis, the number of ketone molecules arising from a fatty acid would

a appointed, the number of actions information a carry and would whole portion of the quotient when the number of carbon atoms in the fatty whose portion of the quotient when the number of caroon atoms in the latty molecule is divided by 4. This bypothesis was adopted by Deucl, Quastel, and morecule is divided by 4. This dypointesis was adopted by Deuti, Quaster, Lelon, Blixenkrone Møller, and Stadie, since it accounted for the greater than 1.1. Account, disconstince assuer, and statue, since it accounted for the greater than 1 1 and of ketogenesis from the higher fatty acids, the lower oxygen consumption hatto of setogenesis from the nigner ratty acros, the ower oxygen consumption than that expected from the 1 1 ratto, and the formation of setone bodies from odd numbered fatty acids (Fig 34)

an numbered larty actus (Fig. 34).

Until recently the multiple alternate oxidation theory was adequate to explain Condition recently the manippe anternate unusation energy was adequate to explain the available data. However, it implied a phenomenon rather difficult to explain the available data stowever, it impures a prenomenou rainer dimetric to expain on block-mical grounds. The simultaneous outdation of every alternate carbon on outcomenical grounds. And simultaneous oxidation or every atternate carbon offered no difficulty. But how could one explain the selective splitting of the acount outsites no auntenty. But now come one explain to eseective splitting of the molecule at every second keto group instead of at every keto group? This difficulty is avoided by a newer conception, which also accounts for other recent evidence to avoided by a newer conception, which also accounts for other recent evidence on compatible with the theory of multiple alternate oxidation. In a systematic in not companies with the theory of multiple attends oxudation. In a systematic in the study of the ketogenic properties of fatty acids consisting of from one to elevemeanusy of the ketogenic properties of fatry series consisting of from one to cieven carbon atoms Jowett and Quastel(1, 12) noted, among other things, ketone pro-

СН, СО СН, СООН СН, СО. СН, СООН СН, СО. СН, СООН СП, СО. СН, СООН	C ₁ H ₁₀ O ₂ + 7O ₂ + 4CH ₂ ·COCH ₂ ·COOH + 4H ₃ O	I mol palmitic acid + 7 mols O1 + 4 mols acetoacetic acid	(No butyric or acetic acid appears at any stage of the reaction)	175 mols Os per mol of acetoacetic acid
CH ₁ ·CC				

Fig. 34 -Hurtley's theory of multiple alternate oxidation (Soskin and Levine [64])

(Palmitic acid)

duction from valence acid (C₂) and a greater production of ketones from hexanoce stational from huttyre (C_i) Since valence and is known to give use to sugar and the man mutyar to some valence and is known to give use to sugar through propionic acid, one can account for the ketone formation only by assum tarough proposite acid, one can account for the actors formation only by assuring a condensation of a two-carbon atom fragment from one molecule of valence IIO use a condensation of a two-carbon atom fragment from one molecule of valence and with a similar two-carbon atom fragment from another molecule. The conactu with a simular two-carbon atom fragments (acetic acid) could also account for censation or such two-carpor; atom histories factor actury come and account for the greater letone formation from hexanosc than from buttyric acid. Leloir and Musioz (21) confirmed the findings of Jowett and Quastel

MacKay and co-workers (42, 43) recently performed feeding experiments on anacsay and co-workers (42, 43) recently performed recoing experiments on intact animals, the results of which supported the interpretation of the above in mact animals, the results of which supported the interpretation of the above in the work and led them to postulate a new theory, which they have termed the sure work and sed them to posturate a new theory, which they have termed the "6 and atton acetic acid condensation hypothesis" They found, in brief, that the P usuation actic acid condensation hypothesis 4 hey found, in thie, that the feeding of propionic acid to their animals led to an accumulation of glycogen in recumg or proprionic acid to their animals ico to an accumulation of gycogen in the liver without formation of ketone bodies. The feeding of valenc acid (C_i) led to both glycogen and ketone body formation. Heptanor, and (C₁) gave rise to to note gyeogen and Ketone poop formation are planned actually gave the to glycogen and to more ketones than did valenc acid. MacKay and associates posgycogen and to more xerous; than one vastic sect. Discreasy and associates pos-tolated that all fatty acid chains whether odd or even numbered, were subjected tuated that an fatty and chains whether out or even numbered, were subjected to oxidation at each alternate carbon atom. However, the molecules then split at to outside at each atternate camou atom stonever, the morecules than sput at every keto group to form a number of acetic acid molecules except where a three every seto group to form a number of actic acid molecules except where a three carbon atom fragment remained to form propionic and (This, of course, resembles in part the original β oxidation theory, although there is little basis for decid ones in part the original p undustion theory, authorize the article class for deciding between successive or simultaneous condition and splitting). Retones are ing our wen successive or simultaneous obtuation and sputting). Actores are formed by the condensation of two molecules of acetic acid (Fig. 35), a process

Friedmann's observation was made on isolated livers perfused with solutions containing acetic acid Recently Barnes et al (45), using acetic acid containing containing acetic acid Recently Dames of at 1451, using acetic acid containing heavy carbon in 11 vilro experiments, conclusively demonstrated this chemical re many carbon in in time experiments, concuratively demonstrated this comment re-action. Weinhouse et al. (46) carned this type of evidence a step further, using action weinhouse et at (40) carried this type of evalence a step further, using containing advoactive carbon in the carboxyl groups octations and outsyris acids containing radinactive cations in the carboxyl groups.

They found that liver slices converted these substances into acetoacetic and pos arey towns that over sites converted these solvatures into accroacetic and possessing radioactivity in the β keto group as well as in the terminal carboxyl group sessing radioactivity in the p seto group as were as in the terminal carooxyl group. This is conclusive evidence that the acetoacetic acid is formed from two-carbon

out tragments
The hypothesis of MacKay and co workers is the most reasonable explanation of the known facts at the present time

For practical purposes the liver may be regarded as the chief, if not the only, source of ketone bodies in the intact organism. The extent to which Letones ac source on Actome Dodles in the IDIACI organism. And extent to which Actories accumulate in the blood or are excreted in the urine will, of course, depend on wheth crating in the most of the extrateroun the name was, or course, depend on whether er they can be disposed of by the extrahepatic fissues and how rapidly such utilizations.

СН1.СО.СН1.СООН	O++ 4HtO	te acid loacetic acid En and Levine [64])
сн, со-сн, соон	C ₁ H ₁₅ Q ₁ + 70, + 8C ₂ H ₁ Q ₂ - + 4CH ₂ ·COCH ₃ ·COOH + 4H ₂ O	I mol palmite acid + 7 mob O ₁ +8 mols acetic acid 8 mols acetic acid
сн, со сн, соон	CaH 204 + 702 + 8C2H	1 mol palmitic acid +- 8 mols acetic acid 1 75 mols O ₂ ; 35
сн, со сн, соон		Fro

СН, СООН СН1, СООН СН2, СООН (Аселс асад)

(Palmitic acid)

KETOSIS 121

tion may occur. Some of the earlier mivestigators regarded the ketone bodies as a hormal intermediary products of fat metabolism, which appeared only when there was a failure in carbohydrate oxidation. It was thought that under these circum stances the ketones could not be metabolized because of the supposed absence of a coupled oxidation phenomenon which ordinarily occurred (47). It is now well recognized that ketosis occurs under conditions in which large amounts of carbohydrate are being oxidized, and, indeed, it has been impossible to demonstrate any relation hetween the degree of ketosis and the rate of carbohydrate oxidation (48, 49, 50, 57). On the other hand, there is ample evidence that both acetoacetic acid and β hydrorybutyric axid are catabolized to CO, and H₂O by kidney, mus cells, heart, brain, testis, etc., as tested on isolated shees m wire (52, 53, 54, 55). Similar evidence is available for perfused whole organs, such as muscles or kidneys (53, 54). The probable pathway of dissimilation of the ketones is indicated in Figure 18 (5) 54).

The rate of utilization of the ketone bodies by the normal intact organism has been estimated hy a number of investigators (55, 56). It is important to note that this utilization, at the blood concentrations of ketones ordinarily found in clinical ketons, may constitute a highly significant portion of the total energy requirements of the organism. Indeed, it has been estimated that ketone utilization in the animals which have heen studied could account for from 50 to 80 per cent of the total oxygen consumption. In view of this great capacity for the utilization of ketones, the small amounts normally found in the blood may indicate that even the normal liver forms, and continues to secrete, some ketone hodies into the blood.

It might be supposed, however, that the severe ketosis of diabetes, phlorhizin poisoning or starvation is the result of some difficulty in the utilization of ketones by the periphery, with or without a greater production by the liver. This possi bility has been tested both in vitro and in vivo, without confirmation Chaikoff and Soskin (14) have shown that the peripheral tissues of the diabetic organism dispose of the ketone bodies as rapidly as do those of the normal animal. This has since been amply confirmed (25 48, 51, 54) With the possible exception of the adrenalectomized animal (68) it must be assumed that, whenever ketones appear in excess in the blood and other tissues this condition is due to a rate of formation and secretion by the liver sufficiently rapid to exceed even the large disposal ca pacity of the periphery. It is thus no longer proper to speak of antiketogenesis in the sense so long employed by clinicians, by which they actually meant ketoly sis (ketone oxidation). In view of present knowledge, the various ketogenic antiketogenic ratios (47) which have been used to calculate the amounts of carbohydrate "necessary for the oxidation of the ketone bodies" must be regarded as being without any real significance

² Crandall and his co workers (57) differ from this opinion on the basis of experiments with the London cannula technic

will be a diminution of ketogenesis—even though some of these substances themselves be ketogene in action if given at a time when the enzyme system unoccupied Such substances are odd numbered fatty acids, critical amino and benzoic, criniamic, and a aminobutyric acids. The type of inhibition they exert is somewhat analogous to the well known action of malonate esuccincidely/drogenesis especting (52).

We may summarize by saying that the ketone bodies are probably normal mediates of fatty acid catabolism in the liver. They appear in excess in the whenever the hepatic metabolism of fat is sufficiently speeded up, either by of carbohydrate substrate or by a disturbance in the normal regulation of the strate mixture. The ketone bodies are readily utilized by the peripheral tunder practically all known conditions. The utilization of ketone bodies may some relationship to the utilization of sugar by the extrahepatic usues, in so these two substrates may compete for the available oxidative mechanism it is evident that the development of a ketosis in the diabetic state cannot ligarded as evidence for the non utilization theory of diabets. It is perhaps compatible with the overproduction theory, for if one broadens the latter cotion to signify the overproduction of metabolic substrates (i.e., sugar plus ket it is clear that the use of the ketones by the peripheral insues will leave a greeness of sucer to accumulate in the blood and shall over into the unite.

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1934

Data Urine nitrogen O, consumption CO, production

0 202 gm/hr 11 105 L/hr 8 200 L/hr

Calculations

1 gm of urme N represents 6 25 gm of metabolized protein

Protein oudized = 0 202 X 6 25 = 1 26 gm/hr

To oxidize 1 gm of protein o 957 L of O. are required and o 774 L of CO, are produced O, used in the oxidation of protein = 1 26 × 0 957 = 1 206 L

and CO, produced in the exidation of protein = 1 26 X 0 774 = 0 975 L

Non protein O. = II 195 - I 206 = 0 080 L and non protein CO. = 8 200 - 0 075 = 7 315 L

Non protein R Q = $\frac{7 \ 315}{0 \ 080} = \frac{0 \ 733}{0 \ 080}$ Percentage of non protein O2 used by CHO ==

cm O₂ used by CHO =

$$100 \left(\frac{0.733 - 0.707}{1.00 - 0.707} \right) = 8.87 \text{ per cent}$$

or CHO oxidation = $\frac{9.98 \times 8.87}{100} = 0.886 \text{ L}$

O used for CHO oxidation

$$O_1$$
 used for CHO oxidation $= \frac{9.959 \times 8.87}{100} = 9.886$

and CO, produced by CHO oudation (RO = 1 00) = 0 886 L O, used for fat oxidation = 9 980 - 0 886 = 0 101 L and CO, produced by fat oxidation = 7 315 - 0 886 = 6 429 L

To oxidize 1 gm of CHO (starch) o 820 L of O2 are required

CHO oxidized =
$$\frac{0.886}{0.820}$$
 = $\frac{1.07 \text{ gm/hr}}{1.07 \text{ gm/hr}}$

To oxidize I gm of fat 2 013 L of O, are required

Fat oxidized =
$$\frac{9 \text{ 103}}{2 \text{ 013}} = \frac{4 \text{ 52 gm/hr}}{4 \text{ 52 gm/hr}}$$

Similar calculations may be made for all levels of the NPRO from 07 to 10 In actual practice, it is customary to ascertain the significance of an R Q de termination by consulting tables or nomograms prepared by Zunz and Schum burg (8), Du Bois (9), and others (5)

THE COMPOSITE NATURE OF THE R O

It is becoming increasingly more evident that the NPRQ of the whole body, like the D N ratio, cannot be regarded as the index of a single process. The ortho dox interpretation of the NPRQ of about o 7 involves the tacit assumption that the only vital processes (aside from protein catabolism) which are in progress and which ultimately consume oxygen and give rise to CO, are those associated with the oxidation of fat Yet there is very satisfactory evidence that other processes which require oxygen or yield CO, are taking place under those conditions. It is generally agreed, for example, that the brain derives its energy solely at the ex pense of carbohydrate and yields an R Q of about 1 o at all times (10, 11, 12 13 14) This high R Q must be balanced by a correspondingly low one in some other tissue or organ if the composite R Q of o 7 obtained from the whole body is to

mean anything at all. Authentic low R Q 's below o 7 have been obtained particularly from the liver, as will be discussed in chapter xm (p 142). It is, therefore obvious that the correct interpretation of an R Q cannot he as simple as that used by its original exponents and some of their present day followers

The conception of constituent R Q 's going to form a composite R Q has actual ly been used to explain values of the R Q over 10. The transformation of carbohydrate into fat, a material with relatively lower oxygen content, would yield a theoretical R O of about R O.

$$4C_4H_{12}O_4 + O_2 \rightarrow C_{16}H_{22}O_2 + 8CO_2 + 8H_2O$$

 $R Q = 7 = 8 O$

This transformation usually occurs when there is a plethora of carbohydrate avail able in the hody. Under these circumstances the R.Q. above unity is said to result from the transformation and the simultaneous oxidation of carbohydrate (5, 7). However, for the sake of convenience, this type of explanation has been confined artificially to R.Q. values over 1.0. It is evident that, if carbohydrate could be converted to fat under conditions where fuels other than carbohydrate were also being oxidized any R.Q. under 1.0 might have a high component due to the transformation, thus abrogating the classical calculations. In reality, there is no evidence that this does not occur. In fact, the work of Schoenheimer and his associates (15, 16), in which heavy isotopes were used as markers, has clearly indicated that there is a constant interconversion of one foodstuff into another even under conditions where no body weight is gained or lost

Catheart and Markowitz (17) and others have shown that the oral administration of 50 gm of glucose to the fasting human causes a leisurely rise in the RQ to values somewhat less than 10, while the administration of equivalent quantities of sucrose, galactose, levulose, or dihydroxyacetone causes a prompt rise in the RQ to values above unity. The more rapid rise in the RQ which occurs with the latter substances cannot be accounted for by their relative rates of absorption from the gastro intestinal tract, and their charmacl composition is theoretically incompatible with an RQ over ro. It is clear, therefore, that even such relatively simple foodstuffs do not yield RQ is which may be reasonably interpreted as re sulting from their oxidation alone

Much has been made of the fact that the RQ of the whole mammalian organ ism has not very often been found to fall below o 7 Indeed, it was formerly ous tomary to ascribe any lower RQ to some undetected fault in technic More recently, admittedly authentic low RQ s have been obtained (18, 10), and other instances in the literature which are similarly free from technical criticisms (19) have been reviewed Some of these low values were obtained in normal human subjects under social conditions of feeding—for example, on high fat intakes before the subjects became acclimatized to the abnormal diet. This is significant because the customary feeding babits of man and of animals have resulted in rather arbitrary conventions as to the number, composition, and size of meals and as to the periods during which RQ measurements of the absorptive and post absorptive states are made. The intake of food is ordinarily spread over a considerable proportion of the 24 hours. This means that all the various oxidations, conversions etc, which yield the highest and lowest components of the composite RQ are usually proceeding simultaneously. Under these circumstances one could hardly expect to obtain anything more than an intermediate range of values for the RQ of the whole body.

To succeed in demonstrating a truer range for the component R Q sof the body on a normal diet, it would be necessary to set the experimental conditions so as to allow the processes responsible for either the lowest or highest component RQ s to predominate temporarily In other words, it would be necessary "to catch the metabolic processes off balance" This has been done by Werthessen (20), who trained rats to eat their entire 24 hour food requirement within a period of 1-5 hours He found that in the same animal, after such a meal, the R O (determined at frequent intervals) varied from extremely low to extremely high values. The range of these variations in all his animals was from 0 27 to 1 70 (See Fig. 37) Markowitz (personal communication), working with Cathcart, performed this experiment upon himself and obtained results similar to those reported by Werthessen These experiments show that the range of RO values ordinarily obtained depends not so much upon the chemical reactions in the body as upon the customary conditions of observation. The extreme RQ values obtained under special conditions again demonstrate that the usual RO s are integrals of higher and lower quotients

The fact that the R Q of the whole body is a composite of many R Q 's originating in different organs and arising from different chemical reactions occurring simultaneously, does not preclude the possibility that all the energy involved may not ultimately be derived from a single foodstuff. When an N P R Q of 0.7 is obtained, it is possible that only fat is being broken down, that some of it is conduced directly in one organ, that in a second organ another portion of the fat is transformed into other metabolites and that these metabolites are oxidized in still a third one. The net result of all these processes could still be an R Q of 0.7. The point is that this figure, by its very nature, depends solely on the starting material and the end products of the series of reactions. It gives no indication whatever of the intermediate reactions. Under these circumstances the characteristic diabetic R Q cannot be interpreted as indicating a lack of ability to ordize carbohydrate. Thus, a fatty acid might break down directly to CO, and H₁O as follows.

The theoretical R Q of this process is 18 - 26 = 0.693 The same fatty acid might first be converted to carbohydrate and then oxidized

$$C_{15}H_{26}O_2 + 8O_2 \rightarrow 3C_6H_{22}O_6 + 18O_2 \rightarrow 18CO_2 + 18H_2O$$

The R Q for this manner of breakdown is also 18 - (18+8) = 0 693

A further characteristic of the diabetic R Q is its failure to rise after the admin istration of carbohydrate as it does in the normal organism. This abnormality may be explained on exactly the same basis as the quantitative excretion of ad

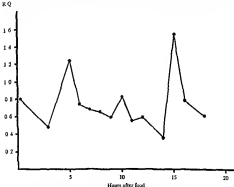


Fig. 37—Serial determinations of the R Q in a trained rat following the intake of its 24 hour food requirement at a single meal (From Werthessen [20])

ministered sugar which we have previously discussed (p 105). It is due to the fact that the extrahepatic tissues of the diabetic organism are already being supplied with a superabundance of sugar so that the administered carbohydrate is not metabolized but overflows into the unine together with the excess arising from the animal so will liver

It is clear that neither the low R Q of diabetes nor the failure of the R Q to nee following the administration of sugar constitute evidence for a fact, of ability to oxidize carbohydrate (For a further discussion of the R Q see chap μn)

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- - matched on of animals

CHAPTER XII

GLUCONEOGENESIS FROM PROTEIN

HE discussion of the D N ratio (chap ix) led to the conclusion that the type of evidence obtained by feeding protein to the depanceatized an mal shows only that some of the sugar which is excreted is derived from the administered protein, and that it is impossible to say to what extent this conversion occurs. When the phlorhizanzed animal is used in the same way, there is the added difficulty of baving to account for a relatively larger sugar excretion than that which occurs in the depanceatized animal.

Somewhat simpler experimental conditions are possible when perfused organs and isolated tissues are used. Since the composition of proteins is variable, the testing of individual amino acids on the isolated organs and tissues is a further sim plification of the problem. The use of amino acids is convenient as regards their addition to perfusates and nutritive media, and the results are quite acceptable as reflecting normal physiology, for both ingested proteins and endogenous proteins are hydrolyzed to amino acids in the intact organism before further estabolism.

The literature up to the year 1930 relating to the conversion of amino acids to carbohydrate was comprehensively reviewed by Rapport (1) Table 15 summarizes the essential information compiled by him and the additional evidence which has accumulated during the intervening years. Data on the conversion of amino acids to β keto acids are also included because of the possible transformation of the later into sugar, a subject to he discussed in the following chapter. The information in Table 15 is derived from the following types of experiments

In atto

- Ammo acids are fed to depancreatized or phlorhizinized dogs and the urine is analyzed for the extra glucose excreted over and above the amounts excreted on previous days
- 2 Amino acids are fed to starving normal animals, and the rise in liver gly cogen is used as an index of transformation to carbohydrate. An increase of the ketone bodies in the blood and unine is taken as evidence of conversion of the amino acids to 8 keto acids.

Perfusion experiments

1 The liver is perfused with blood to which the various amino acids are added A rise in the glucose or ketone content of the perfusing blood is taken as evidence for transformation

TABLE 15*

AVAILABLE EVIDENCE FOR GLUCONEOGENESIS AND KETOGENESIS FROM THE AMINO ACIDS 1

		In the Experiments			Perfusion and se suise Experiments					
Амино Асца	To Carbo hy- drates	Ke-	References and Remarks		To Car- bohy- drates		References and Remarks			
Glycine	+00+0	0	Lusk (14), phlorhamized dog Pflueger (15), normal dogs Wilson (16), normal rats Butts (17), normal rats MacKay (12), normal rats Olsen (11), normal rats (150 tope carbon as tracer)		0		Bach (5), liver and kidney perfusions and shees Bach (6), liver shees			
Alanine	##	000	Lusk (14), phlorhizmized dog Butts (17), normal rats Wilson (16), normal rats		‡		Embden (18), liver perfusion Krebs (7), liver shees			
Senne	+	0	Rapport (1), phlorhizmized dogs Butts (17), normal rats Dakin (20), phlorhizmized		+		Chargaff (19), liver extracts			
Valine	 	1 1	dogs Butts (21), normal rats Rose (22) phlorhizmized dogs	1	Ì	1				
Leucine	{ :	+	Butts (23), normal rats Dakin (20), phlorhizmized dogs	'	٠	#	Embden (24), liver perfusion Edson (13), liver slices			
Isoleucine	∦ ‡	+	Butts (23), normal rats Dakin (25), phlorhimized dogs	ļ,		1				
Norleucine	: +	:	Butts (23), normal rats Lusk (14), phlorhamized dogs	۱,	. 1	- 1	Krebs (7), liver and kidney			
Aspartic	K 💷		Butts (26), permal rats	١٦	1	- 1				
Glutamic	{		Lusk (14), phlorhizmized dogs Wilson (16), normal rats Butts (26), normal rats	1	1		Weil Malberbe (27), liver			
Argunne	[i + j	۰	Dakın (20), phlorbizinized		П	- 1				
Ornsthine	lo++		Butts (28), normal rats Dakin (20), phlorhizmized dogs			-				
Lysine		0	Dakin (25), phlorhizmized dogs Butts (25) normal rats		1	1				
Cysteine	+		Dakin (25), phlorhizmized dogs	+		- 1	imythe (29), liver slices			
Cystine Methionine	0	- 1	Butts (30), normal rats Transformed to cystine (q v) Dakin (20), phlorhizmized	+	1		mythe (29), liver slices Embden (24), liver perfusion			
Phenylalanine	+	. (dogs Butts (31, 32), normal rats		1	2 E	ldson (13), hver shees			
Tyrosine	(0-1 +	+ 1	Lusk (14), phlorhizmized dogs Butts (31, 32), normal rats			LIF	imbden (24), liver perfusion idson (13), liver sices			
	10		Dakın (25), phlorhızınızed dogs							
Histidine	{+}	0	Remmert (33) and Feather stone (34), normal rats							

^{*} Zero indicates negative experimental results indicates no data

TABLE 15-Continued

			In the Experiences	Preparion and in mile Experiments			
Амию Астр	To Carbo- by- drates	To Xe- tones	References and Remarks	To Car- boby- drates	To Xe- tence	References and Remarks	
Tryptophane	0	0	Dakin (25), phlorhazmized dogs Borchers (35), normal rats				
Proline	‡	0	Dakin (25) and Kapfhammer (36), phlorhiz inized does		۰	Edson (13), liver slices	
Hydroxyproline	` +	٥	Kapfhammer (36), phlorhaz inized dogs		+	Edson (13), liver slices	

In vitro

r. Tissue slices (generally liver) are incubated in the Warhung respirometer with various amino acids, and the rise in total carbohydrate, carbohydrate intermediates, and ketone body content of the slices is measured.

2 Enzyme preparations from animal tissues are employed to follow the path-

way of the intermediate metabolism of amino acids

It may be seen that a large part of the evidence collected in Table 15 was on tained in 810, using the D N ratio or the increase in liver glycogen content as the cintemo for carbohydrate formation. The same objections as were raised against the use of the D N ratio in the study of gluconeogenesis from protein also apply in the present connection. The increase in liver glycogen after amino acid admin istration was not regarded as a quantitative index, even by those who used this criterion. This leaves the perfusion and the sin sin c experiments as the possible ource of reliable quantitative information. When all the quantitative vedicace is summarized, it may be seen that definite information is available about only six amino acids. Alanine, aspartic acid, and glutanuc acid are converted to carbohydrate in definite proportions and by known pathways, as follows

Lysine, tryptophane, and leucine are not converted to any measurable degree.

There our quantitative information ends

This leaves fifteen amino acids about which only qualitative information is available, and the information we do have casts considerable doubt upon the validity of even this type of conclusion For example, the in time evidence as to gluconeogenesis from glycine is contradictory, only two out of seven sets of experimenters having obtained apparently unequivocal evidence that this occurred The in vito evidence as to the metabolic fate of glycine is not wholly clear, and it is contradictory in some respects. It is well established that glycine is one of the building stones of creatine (2, 3, 4) and that it may condense with a letocods probably forming new amino acids (5). However, there is no unanimity of opinion as to the deamination of glycine. Thus, Bach (5, 6) found that neither ladney nor liver slices were able to deaminate glycine. Moreover, the standard amino acid oxidase preparations exert no effect upon this amino acid (7). However, very recently Green et al. (8) prepared a glycine oxidase system from ladney which converts glycine to glycovite acid.

CH₄NH₄ COOH + NH₄

Another enzyme system converts glyoxylic acid to oxalic acid (COOH COOH) (8), but, since previous work has shown that oxalic acid is not further convertible in the animal body (9, 10), the work of Green indicates that glycine does not by it self give rise to glucose

This conclusion is strengthened by the work of Olsen et al. (11), who fed so-topic glycine to rats. The liver glycogen showed a delayed rise (confirming Mac Kay [12]), but this glycogen was not derived from the administered glycine, for it did not contain any of the heavy carbon. Olsen et al. (11) drew the important conclusion that evidence concerning the conversion of amino acids to glucose derived from in vivo and in vitro experiments should be re-examined, using labeled amino acids. It is not sufficient to show extra glucose exerction or increased liver glycogen. To be unequivocal, the evidence must show that the newly formed glucose or glycogen is built up from the constituent atoms of the amino acid under investigation.

To cite another example, prohie administered to phlorhizinized dogs has been to cite another example, prohie administered to phlorhizinized dogs has been to cite another example, prohie administered to phlorhizinized dogs has been to cite another example, prohie administered to phlorhizinized dogs has been to cite another example, prohie administered to phlorhizinized dogs has been to cite another example, prohie administered to phlorhizinized dogs has been to cite another example, prohie administered to phlorhizinized dogs has been to cite another example, prohie administered to phlorhizinized dogs has been to cite another example, prohie administered to phlorhizinized dogs has been to cite another example, prohie administered to phlorhizinized dogs has been to cite another example.

We may summarize the present knowledge by saying that, whatever its empirical usefulness, the figure of 44-58 per cent commonly used in metabolic and nutritional work to calculate the carbohydrate equivalent of protein has no real bass in fact. Even under the simplest conditions, using amino acids and the in time technic, it has thus far been possible to ascertain the quantitative fate of only a few of the amino acids. It is evident that much work remains to be done in this field

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CARBOHYDRATE METABOLISM

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tomy experiments for the sugar utilization of the extrahepatic tissues of normal and depancreatized animals, calculated that sugar must be formed from fatty acids in the livers of both types of animal, as follows (a)

hour, ie 60 gm per day for a 10 Kg dog The average artenovenous blood ditterence in fasting animals is 4 mg per cent [Cori (5)] and if the tissues of a fasting 10 Kg dog are absorbing sugar hand a series of the about 100 ce per 10 Kg dog is a stully

and fall sham annum. Foh may and the other could be a Nordon like

or so of fasting the source of 60 - (25 + 6 + 6) = 23 gm of the daily hepatic sugar production of the 10 kg dog is unaccounted for The discrepancy is 50 gmat that it seems impossible to account for the facts without assuming considerable conversion of fatty acid to sugar in the liver of the facting dog

Departreatited dog - Mann's [11] observation that hepatectomy of a previously departrea and dog regulated in a fall of blood sugar level similar to that occurring in a normal hepatec

producing at least 45 gm of sugar per diem for a 10 M, use two at the use to not sugar per diem of sugar and a-c sm of introgen per diem. The total sugar pro-

sugar a mani

and II will 17 and not not had

DIRECT EVIDENCE

23) Our present knowledge of tissue-enzyme chemistry and of intermediary fluctuo

olsm indicates the existence of suitable pathways for gluconeogenesis from fatty acids (chap iu, p 54). The point at issue, therefore, is not whether the process can occur but whether it does occur in the mammalian organism.

In view of Young's calculations, it is of interest to consider why the administra tion of fat to experimentally diabetic animals has usually not resulted in sufficient excretion of extra sugar to indicate gluconeogenesis from fatty acids when the cal culations were made on the basis of the classical interpretations of the D N ratio (24) This is not surprising when it is remembered that these interpretations, by their very nature, practically exclude the possibility that such calculations might yield positive results. Even so, it might still be possible to show extra sugar excretion if the experimental animal could make additional amounts of sugar over and above that which it is already forming from endogenous protein and fat, including the amount which is being utilized during the experiment But this involves the unwarranted assumption that the capacity of the liver for gluconeogenesis from fat has not been reached before the fat is administered. The fact that this is not the case for protein has no hearing for it happens that fat is the only stored food suhstance present in practically unlimited amounts so far as the daily requirement of the hody is concerned It might therefore be expected that fat would be used to capacity when the liver is forming sugar at an uncontrolled rate

From the practical standpoint the experimental procedure to test the extra sugar excretion involves the administration of fat to the diabetic animal on the fourth or fifth day after panerastectomy after the withdrawal of insulin, or after starting phlorhizination. At this time the animal is suffering from acute diabetes with ketosis, and the administered fat makes him even more sick. In certain experiments in which some extra exerction of sugar after fat administration was reported (25), the animals died shortly. In order to obtain positive results by this method, it is apparently necessary to exceed physiological limitations to a degree incompatible with life.

There have been a number of experiments the results of which favor glucones genesis from fat even though the investigators did not take into account the factor of utilization in these experiments neutral fat or fatty acids were administered to intact normal or diabetic animals, or certain hormones (e.g., epinephini) or drugs (e.g., pilothinin) were given to such animals in an attempt to force excessive gluconeogenesis from endogenous fat stores. The results of these experiments were judged by the increases in carbohydrate content of the liver and muscles of the normal animals and by the increased sugar exerction of the diabetic animals. As might be predicted from our previous discussions of the dynamic balance and the D N ratio, these experiments have yielded both positive (3, 25, 26, 27, 28, 29, 30, 31) and negative (24, 32, 32) results Under the circumstances it is justifiable to place greater weight on the positive than on the negative findings. This evidence and preceding work of a similar kind have been comprehensively reviewed by

Macleod (3) and Geelmuyden (2) and will not be discussed here. It will be more profitable to confine the discussion to more recent and less controversial evidence

The theoretical R Q for the cooversion of protein to carbohydrate has been variously calculated as o 613 (Magnus Levy [34]), o 632 (Lusk [35]), and o 706 (Geelmuyden [36]) The R Q for gluconeogenesis from fat has been calculated to be about 0.28 by Pembrey (37) and by Macleod (3) The theoretical R Q for ketogenesis from fat may be calculated to range from 0.65 to 0.00, depending upon the number of molecules of β hydroxyhutync acid which are supposed to anse from one molecule of fatty acid. The work of Bluxcokrone-Myller (38) strongly indicates that the value hes closer to zero than to the higher figure

Since gluconeogenesis and ketogenesis occur primarily in the liver, it would be expected that RQ determinations performed on the isolated liver under the appropriate physiological conditions should yield very low values. This is the case Gemmill and Holmes (39) found that the RQ of liver slices from a rat fed on a normal det averaged o 79 while that from a rat fed hutter averaged o 58 Stade and co-workers (40) observed RQ 's of about o 32 in liver slices from the depan creatized cat Similarly in the perfused livers of normal and depancreatized cats Bluxenkrone Møller (38) ohtained RQ values which averaged o 57 for the normal

coneogenesis from fat. But the simultaneous occurrence of gluconeogenesis illumination, and particularly of variable ketogenesis, makes it difficult to use the RQ as a quantitative index. Evidence based upon chemical determination of newly formed carhohydrate or carbohydrate intermediates is more convuncing

We have already mentioned the work of Gemmill and Holmes (39), in which they found very low R Q values in the isolated liver slices of hutter fed rats. They also observed a coincident iocrease in the carhohydrate content of these slices which was greater than the iocrease observed in liver slices taken from rats on a

each substance and in practically all tissues they observed a large production of lactic acid. The simultaneous decrease to the carbohydrate content of the tissue when it occurred, was significantly less than the iocrease in lactic acid. In the case of the liver, when oxygen was present there was an increase in the carbohydrate content as well as in the amount of lactic acid. It was obvious that the lactic acid could not be accounted for as ansing from carbohydrate. The authors considered the possibility that the added fatty acids might have stimulated the production of

lactic acid from some other substance but they concluded that this supposition nature actor from some other sumstance out they constituted to at this supposition could not be justified. They pointed out that in the brain and liver for example come not be Justiness 1 new pointers out that in the orain and over 100 example they were dealing with tissues which ordinarily produce hitle or no lactic acid and taey were deating with ussues which ordinarily produce attic or no factic acid and which contain no other known precursor of factic acid. Their work therefore 143 which contain no other known precursor or factic acts. Their work inferiore yields convincing evidence for the formation of carbohydrate from fat through a yieus convincing evidence for the formation of cationydiate from fat introduce lactic acid stage (probably via pyruvate) More recently gluconeogenesis from sactic acid stage (propanty via pyruvate) ander recently gracouleugenesis atom fat in isolated mammalian tissue has again been confirmed by Weil Malherbe (43) nas in issuateu mannaisin tissue nas again peen communeu by wen prainterpe (43) who demonstrated the *in who* formation of sugar from added accloacetic acid by

Another method by which the extrahepatic utilization of sugar has been ex Amounter method by wanth the extranspatic utilization of sugar has over a cluded and one which is a step nearer the intact organism is the perfusion of the enace, and one which is a step nearer the intact organism is the periusion of the isolated whole liver. This method is not easy, and it is sometimes difficult to obtain usuated whose liver 1 his method is not easy and it is sometimes diment to obtain satisfactory preparations (44) Nevertheless a number of competent investigators sausactory preparations (44) Nevertueiess a number of competent investigations are carried out reasonably successful liver perfusions as judged by a maintained rate of flow of perfusale through the hver with httle or no edema the continued auce or more unperturante uniougn the over with name or no evental the continued excretion of hile and the storage of glycogen. Burn and Marks (45) perfused the exercision on one and the storage of giveogen, but in and marks (45) perfused the glycogen poor hyers of fat fed dogs and of a depancreatized cat. A large production gyvogen poor uvers of fat fed dogs and of a depancreatized cat. A large production of actione bodies and of sugar was observed. The pre-existing carbobydrate con ou actione boules and of sugar was observed. And pre-existing carbodydrate content of the livers accounted for but a small fraction of the sugar which appeared tent of the livers accounted for but a small fraction of the sugar which appeared. The disappearance of lactic acid was ruled out as a factor. As regards gluconeoane disappearance of factic acid was fuscif out as a factor. As regards gluconeo-genesis fum protein. Burn and Marks rightly (in view of our previous discussion general num protein thurn and matrix nightly un view of our previous discussion of the D N) rejected the use of any of the orthodox values for the D N ratio In or the DAN rejected the use of any of the orthodox values for the DAN ratio in stead, they calculated that if all the carbon in the protein molecule were recomaread they calculated to at 11 an the carbon in the printern molecule were recom-blined so as to form dextrose the ratio of dextrose produced to nitrogen set free in oned so as to form dextrose the ratio of dextrose produced to introgen set tree in the form of trea and ammonia cannot be greater than § 3.1. Values for the $D\ N$ the form of the and ammonia cannot be greater than 0.3.1 values for the D N ratio above this figure would therefore demonstrate gluconeogenesis from fatty ratio above tins figure would inference demonstrate guiconeogenesis from fatty and Out of a total of forty seven determinations of the D. N ratio thirty two exceeded the value of 8 3 and in seven cases the ratto tose above 17 0 reueu tae vatue of 3 3 and in seven cases the ratio rose above 17 o Heller devised ingenious methods (40) to observe the sugar output of the liver

retuer devised ingenious methods (40) to observe the sugar output of the uven in the in normal and phlorhizinized cats anesthetized with Pernocton After de 33 лиц п поппы and pniornizinized cats ancidetized with remotion After de ducting the amounts of carbohydrate which might have come from glycogen fac uncing the amounts of carbonydrate which might have come from give ogen it acid and giverol be calculated D N ratios ranging from $5 \circ 10 \cdot 18 \circ (47)$

More complete and conclusive work upon the subject was done by Elexenkrone Addiet (48) He perfused the livers of normal and of phlorhizmized cats with sodi aspure (45). He periused the avers of normal and of paramazanzed cats with soci um butyrate. After accounting for other possible sources of carbohydrate, he ob our outgrate After accounting for other possible sources of carbohydrate he ob-lained D N ratios ranging from 10 o to 20 or over Perfusion with sodium suctance U , V ratios ranging from 10 o to 20 o or over recreasion with social successful V ratios as high as 42 o. He concluded that about 20 per cent of cuate yielded D N ratios as figures as 42 \circ 11 \circ concluded that about 20 per cent of the added butyric acid was converted into kelone bodies and that the remainder are added butyric acid was converted into ketone bodies and that the remainder went to sugar via succinic acid. Cat hvers were perfused with blood according to a technic worked out by the author In control experiments this technic per ecume worked out my the author in control experiments has recome per mitted glycogen storage from glucose etc, thus demonstrating preservation of

normal hver function Chemical determinations included glycogen, fat, and ke tone content of the liver before and after the perfusion, blood sugar, ketones lac tic acid, urea, oxygen, and CO, at frequent intervals. Sodium butyrate was added to the perfusing blood after a control period. Table 16 shows a typical experiment performed on a liver from a normal cat (48).

It can be seen from Table 16 that the carbobydrate, newly formed in a liver per fused with sodium butyrate, could not have arisen from protein conversion and must have been derived from the fatty acid Unequivocal confirmation of this conversion was supplied by Hastings and co workers (49), who fed butyric acid containing "heavy" C atoms to normal rats and found the labeled C in the liver give

TABLE 16

PERFUSION OF NORMAL CAT LIVER WITH SODIUM BUTYRATE (BLIXENEROVE MØLLER [45])

Liver weight, 61 mm blood volume 200 cc sodium historiae added 200 mm

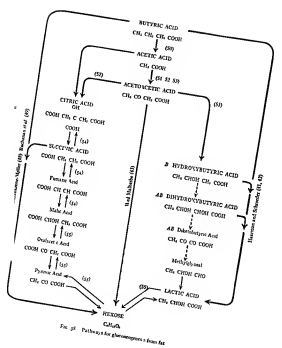
	Соиски	RATION (BIG	PER CENT)	AMOUNT	Kinvere				
CHEMICAL ANALYSIS	Intal	Fael	Difference	(Mg)					
Total ketones of blood Total ketones of liver	39 3 33 0	93 5 76 0	54 2 43 0	162 6 26 2	#88 8 mg of ketones appeared				
Blood sugar Liver glycogen	322 0 80 0	304 0 235 0	82 0 155 0	246 0 95 0	341 0 mg of earbohydrate appeared				
Blood ures	58 o	700	120	35 o	Corresponds to the breakdown of 106 o mg of protein				

In control experiments with butyrate 75 mg of ketones and 54 mg of carbohydrate were formed. The breakdown of protein could have given rise at the utmost to roo rag of sugar. The biliance research therefore that the sodium butyrate gave rise to—

cogen of these animals. These findings are quite in accord with other pertinent evidence discussed elsewhere in this volume. We have seen that fatty acids are broken down to the ketone bodies by way of acetic acid (chap x). Acctoacetic acid may condense with oxalacetic acid to enter the tricarboxylic acid cycle, which is the common reservoir for derivatives of all three major foodstuffs (56). And each mem ber of the cycle has been shown to be capable of resynthesis to glucose (55). In addition, acetic acid may, under certain circumstances, enter directly into the tricarboxylic acid cycle without going through the acetoacetic acid stage (52) (see chap mi, p. 54).

Figure 38 graphically summarizes the more direct evidence for gluconeogenesis from fat and indicates the intermediate chemical steps by which it may occur We may conclude that this process can and does play an important role in both the

normal and the diabetic mammahan organism



CHAPTER XIV

UTILIZATION, DISSIMILATION, AND OXIDATION OF CARBOHYDRATE

HE use of the term "oxidation" to describe the complete breakdown of a foodstuff to CO, and H_iO in the tissues carries with it certain traditional physiologic connotations which are no longer acceptable in the light of present-day biochemistry. Chief among these is the old conception that the ong nal foodstuff can liberate its energy for use by the tissue by the simple addition of orgen to its atoms. But, as was shown in chapters ii, iii, and iii, the oxidative breakdown of the energy materials in the tissues is actually a far more complicated matter, involving the processes of oxidoreduction, decarboxylation, addition of CO, phosphorylation, hydrolysis and transamination.

It is true that the net result of a whole series of reactions may be written as if it were a simple oxidation, as, for example

$C_4H_{12}O_4 + 6O_2 \rightarrow 6CO_2 + 6H_2O$

Indeed, it was our limited knowledge of the intermediate steps in this equation which originally led to the inaccurate use of the term "exidation" But, now that most of the intermediate steps are known the continued use of "oxidation" for the allower process is a source of great confusion. For example, when the buchem ist speaks of the "oxidation of lactate," he means specifically the withdrawal of hydrogen from lactate with the formation of pyruvate. The physiologist uses the same words to denote the breakdown of lactic and to CO, and H.O. It would be far better for all branches of biological science to use the term "oxidation" in its strict chemical sense, and this is the sense in which it is used in this volume. For the complete breakdown of a substrate to CO, and H.O. we employ the term "complete oxidation" or "dissimilation" (1)

There is a practical need arising out of the conditions of experimental work for another term, namely, "utilization". In working with the whole living organism or even with isolated tissue or time, it is office possible to follow the disappearance of a substrate from the blood or nutritive medium or from the tissues themselves without being able to ascertain the extent to which the oxygen consumed and the CO, evolved in the interim were actually concerned with the substrate that disappeared. Other substrates are necessarily always present under these combineds and their participation in the reactions under observation is not necessarily ruled out by an approximate equivalence between the respiratory exchange and the dis

appearance of the experimental substrate. Such equivalence may be coincidental, for it also happens, not infrequently, that the disappearance of a substrate bear no discernible relationship to the respiratory exchange (2, 3). Under these circum stances when it is impossible to determine the exact chemical fate of the substrate which is disappearing, it is best to employ the term "utilization". As used in this volume, and applied to carbohydrate, for example, it means the disappearance of sugar from the blood or nutritive medium or tissue without storage as glycogen or activalishing as herose or later early

UTILIZATION OF CARBOHYDRATE AS DETERMINED BY THE DISAPPEARANCE
OF THE BLOOD SUGAR IN LIVERLESS ANIMALS

The rapid disappearance of the blood sugar after removal of the liver from the normal animal has been discussed in chapter vii, in connection with the site of for mation of the blood sugar. The mere withdrawal of sugar from the blood by the extrahepatic tissues cannot, of course, be regarded as proof of its utilization by those insues However, it has been the universal expensione that the carbohydrate content of the tissues and the accumulation of factic acid or any other substance in the blood do not account for the sugar that disappears from the blood of the liveriess animal (4 5 6). The rate of disappearance of blood sugar in such animals may therefore be taken as at least a rough indication of the utilization of sugar by the extrahepatic tissues.

In view of this it is significant that the bfood sugar disappears after hepatectomy or abdominal evisceration in animals which have been supposed to have coased utilizing carbohydrate, as judged by the D N ketosis and RQ exhibited before removal of the liver. Such evidence is available after hepatectomy of depan recatized birds (7), dogs (8), and rabbits (9) and after evisceration of phlorbizinized dogs (10), of depancreatized and pituitary-diabetic dogs (11), and of normal dogs fasted to the point of so-called "hunger diabetes" (12). A similar incongruity between the conclusions drawn from the classic metabolic criteria and the disap pearance of the blood sugar occurs after hypophysectomy of the depancreatized dog (13, 14) and during prolonged injections of epinephrin in the normal dog (15, 16).

UTILIZATION OF CARBOHYDRATE AS DETERMINED BY CHEMICAL BALANCE STUDIES IN LIVERLESS ANIMALS

The groundwork for future chemical balance studies of carbohydrate utilization was laid in the laboratory of H H Dale At that time practical methods for total abdonunal evisceration in the cat were not available. The liver was left in situ with its afferent blood supply tied off. However, the asphyraated organ (with a high free sugar content) could still contribute sugar to the blood by seepage into the vena cava. In their later experiments Dale and his co workers (4, 17, 18) recontribute sugar to the supplements of the vena cava.

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Tinder the m t = 2
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consumes of from the CO produced II a stable intermediate substance of known chemical composition is formed the R Q may be used to esleuiste the course of the reaction (co) Hower it is usually also necessary to determine the amount of original substrate which has disappeared or the amount of intermediate substance.

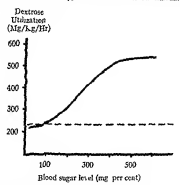


Fig. 39 —The relationsh p between the blood sugar level and sugar ut) zat on in existerated normal dogs (Sork n and Lev ne [5])

which has appeared by chemical analysis. When a single substrate is acted upon by an enzyme system and an unknown stable intermediate substance is formed the difference between the theoretical R Q for the complete orndation of the substrate and the actual R Q obtained may suggest the probable identity of the unknown intermediate (20)

There is no tissue which does not contain a number of substrates and more than one easy me system In working with a tissue it is therefore desirable to allow it to approach the minimum level of autorespiration (i.e. to exhaust its own substrates) before the substrate under investigation is added. If the RQ of the subsequent

reaction agrees with the chemical determination of the disappearance of the added reaction agrees with the chemical determination of the disappearance of the added substrate and the appearance of end products it may then be concluded that the substitute and the appearance of ent products it may then be continued that the particular enzyme system which it was hoped to engage has operated and that the supposed course of the oudstive process has been confirmed

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uppersecutions of the orogative process has been committeed. It is thus apparent that even when one can control the other activities of an It is thus apparent that even when one can control the other activities of a soldted tissue and is dealing with a single substrate the R Q is merely confirma isotated tissue and is usuamy with a single sourcease. The A.V. is metery committee to the information obtained by chemical analysis. When used alone the R.Q. ony to the miorination obtained by chemical analysis. When used alone the K.Q. can at most merely suggest the probable pathway of a reaction, which must then can at most merely suggest the probable pathway of a reaction which must men be demonstrated by chemical means. To illustrate the lack of preciseness of the in occurrous states by cocumeat means as mustrate the tack of preciseness of the in-dications derived from the R.O. Let us suppose that the substrate is hexose and occasions derived from the K Q set as suppose that the substrate is nexose and that no other foodstuff is involved. Let us simplify matters further by considering the possible pathways open to just one of its important intermediary metabolites namely pyruvic acid

ancy pyruvic acio Table 17 summarizes the rather formidable list of possibilities with the experi same a summarizes the variet interaction observed total R Q s for pyrivic acid which are cited have been obtained in different tissues and under different cit ecum name are circu have been obtained in outerent ussues and under different crimination of the individual reac tions favored by the experimental conditions. It is obvious that the total R Q of tunn tayored by the experimental conditions at is obvious that the total $\kappa \ V$ of a single tissue, like that of the whole body is a composite of many possible R Q 3 a surger unsure time that of the whose usely is a compensate of many possible $\kappa \ V$ s. It is also clear that to gain more than the vaguest indication of the fate of the suba as asso ciear unat to gain more main the vaguess muccation of the rate of the sun state from the R Q alone is a mathematical impossibility. Furthermore, when the state from the a Q stone is a mathematical impossibility. Furthermore when the chemical determinations have been made, there is little information that the total R.Q. can add except to act as a check on the possibility that one or more of the end products might have been missed

If we now attempt to apply the foregoing to the interpretation of the R Q in one turner is one further computation which must be mentioned in the body the three main foodstuffs or their breakdown products are constantly available and may be metabolizing simultaneously. It has been shown that amino acids may way to meranomang sumuntancousty it has occu shown that amino acids may yield the same R Q of unity as is given by carbohydrate (21) Acetoacetic acid if reactive same $\alpha \vee \alpha$ in that was a given by calloury trace $\alpha \vee \beta$ according and also yield an $R \circ Q$ of $\alpha \circ \beta$ in view of the limited significant of the second state of the second state of the second cance of the RQ of a single tissue acting on a single substrate what possible mean tancou the κ V of a single insue acting on a single substitute what possible mean ing can be assigned to the composite R Q derived from many tissues acting on a

In this predicament the proponents of the R Q have sometimes resorted to the argument that when the R Q of the whole body is determined over a sufficiently one period of time at must represent the resultant of all the R Q s in all the tissues only period of time it must represent the resultant of an time $K \setminus V$ s.in an time tissues and must therefore ultimately depend upon the chemical composition of the original configuration of the original configurat and must ineterore unimatery depend upon the enemical composition of the original substrates being outlized. This ignores (a) the fact that what constitutes a and substitutes being origized anis ignores (a) the fact that what constitutes a sufficiently long period of time under various conditions is difficult to determine on ventury one period of time under various conditions is anacent to determine (in any case practical reasons have usually dectated rather short periods of R Qwe say case practical reasons have usually dictated father short periods of KQ measurement in the past [22–23–24]) (b) the possibility of partial decarborylation

of some of the intermediary metabolites of the original substrate without further oxidation of the residues, so that the integral of the RQ 's could never equal the theoretical RQ of the original substrate, (c) the possibility that some oxygen is used in the formation of storage or excretion products without the formation of equivalent amounts of CO, with the same result as in (b), and (d) the recently discovered mechanism whereby CO, hitherto considered to be immediately and

TABLE 17

Lyperimental and Theoretical R Q's for the Reactions of Pyruvic Acid (Soskin [ad])

REACTION PRODUCTS	PER	MOL FOL	Tutoberical	Reyysences						
KAACHON PRODUCTS	Con stamed	COs Fro- duced	RQ							
CH, CHNH, COOH (Alaune)	۰	۰	°-°	Braunstein and Kritzman (37)						
C4H12O6 (Hexone)	-0 S	••	-0 5 =0	Benoy and Ell oft (38)						
CO'+H'0	2 S	30	1 = 1 2 2 S	Long (20)						
COOH • CH,CH, • COOH (Succinic acid)	0 75	1 0	1 0 75 = 1 33	Ethott and Greig (39) Weil Mal herbe (40) Rrebs and Johnson (41)						
CH ₄ • COCH ₂ • COOH (Acetoacetic acid)	0.5	10	05 =2	Krebs and Johnson (41, 42)						
CH _s · COOH+CO _s (Acetic zcid)	0.5	10	1 = 2 0 S	Long (20)						
CH ₃ • COOH + CH ₃ • CHOH • COOH +CO ₂ (Acetic acid) (Lactic acid)	0	0.5	<u>0</u> ≤ «	Krebs and Johnson (41)						

OBSERVED R Q & OF PYRUVATE IN VARIOUS TISSUES

	Tassie	Observed R Q	References					
Liver kidney Testis Brain Brain Liver		a 82-t it 1 07-1 24 1 17-t 41 1 18 1 28 t 28 1 19-t 76	Bach and Holmes (43) Elhott and Schroeder (44) Elhott, Greig, and Benoy (45) Elhott, Greig, and Benoy (45) Long (20) Elhott Greig, and Benoy (45)					

quantitatively excreted, may be held hack (temporarily, at least) and its carbon used for the synthesis of metabolic intermediates (25, 26, 27) For example

It is possible that under special experimental conditions, such as prolonged fast ing or exclusive high earbohydrate feeding, the R Q does depend largely upon the chemical composition of the original food material which is being dissimilated But even if this possibility be granted, it is perfectly clear that the composite R Q cannot be used to judge the intermediate steps undergone by the foodstuff on its way to complete degradation to CO, and HiO. In other words, even if we suppose that the R Q of o 7 means that the animal is living at the ultimate expense of fat, there is no reason for the further supposition that the fat is being directly and completely andized in the extrahepatic tissues (see chap m). Thus, the R Q has no weight against the previously cited direct chemical evidence that, in its ultilization, fat is converted to hexose and ketones by the liver and that these intermediates are dissimilated by the extrahepatic tissues.

ATTEMPTS TO VERIFY R Q BY SIMULTANEOUS DETERMINATION OF CARBOHYDRATE UTILIZATION IN INTACT ANIMALS AND IN ISOLATED TISSUES

Despite the inherent limitations of the R Q, a number of investigators have been direct evidence of its validity as a quantitative index of the type of food stuff that is being dissimilated. These attempts have usually consisted of a quantitative comparison of carbohydrate dissimilation as calculated from the R Q, with carbohydrate utilization as determined by chemical balance studies (2, 3, 4, 18 28, 29, 30).

In view of the distinction that we have drawn between dissimilation and utilize tion, it is evident that they need not tally even if the R Q were a reliable index of complete oxidation, for it would be quite possible for more carbohydrate to be utilized than was dissimilated if some of the carbohydrate were simultaneously being converted into fat or another stable form. There is still another difficulty when such comparisons are attempted in intact animals. It has been pointed out (chap vii) that the blood sugar level represents a dynamic balance between the rate at which sugar is entering the blood stream from the liver and from any exogenous source and the rate at which it is being removed from the blood by the tissues of the body. Thus, a rise in the blood sugar level may result either from an increased rate of sugar supply or from a decreased rate of sugar valuation, or from both together. Conversely, a fall in the blood sugar level may be due to decreased supply or increased utilization, or both Nor is it possible to tell which factor is responsible from the mere change in blood sugarlevel unless one is controlled or deliminated

while the other is observed. It is, therefore, futile to attempt to determine the amount of carbohydrate which has been utilized by an intact animal by estimating the difference between its total carbohydrate content at the beginning of an experimental period (plus any sugar which may have been administered) and its total carbohydrate content at the end of the period, for in this procedure the amount of carbohydrate being supplied by the liver is unknown, and any effected sugar administration on this supply cannot be estimated. The experimental conditions are simpler in liverless animals or in isolated tissues, where the available car bohydrate can be estimated or controlled by the investigator

Table 18 summarizes the data of all papers available to the authors from which a comparison of utilization, as determined by chemical balance, and of supposed dissimilation as judged from the RQ, may be attempted A study of the table obviates the necessity for much discussion. It is clear that in eviscerated animals and in isolated tissues, as well as in intact animals, there is no correlation between the results of chemical balance studies and R O calculations. In view of the frequency and extent of the discrepancies, the few instances in which the results hap pen to coincide may be regarded as purely fortutous. A somewhat better correla tion is obtained in isolated brain tissue than in isolated muscle of the whole living animal This may be ascribed to the fact that the highly specialized nervous tissue does not possess the ability of other tissues for storage and interconversions of foodstuffs and, so far as we know, derives its energy solely from carbohydrate (31, 32, 33, 34) (see chap 1 p 16) However, even under these circumstances, the cor relation between chemical balance and the RO is by no means good. This is 10 even in experiments in which the present authors have improved on the usual tech nic of chemical balance by a rapid freezing of the control samples (Table 18 no 9)

As was discussed earlier in this chapter, the blood sugar level has an important influence on the utilization of carbohydrate by the living organism Germill (1) showed a similar influence of the concentration of sugar in the medium on the car bohydrate utilization of isolated muscle in vitro. The various data in Table 18 lack a certain amount of comparability because the other investigators fulled to take this factor into account. Figures 40 and 4 graphically summarize the work of Germill (3) and hitherto unpublished data of the present authors for the evis cerated dog isolated muscle, and isolated brain tissue, respectively, in which car bohydrate utilization and R.Q. calculations are considered in relation to glucose contentration. It is apparent that except for isolated brain tissue, there is no concentration of glucose at which carbobydrate utilization and R.Q. calculations co-incide.

One must conclude that chemical balance experiments offer neither theoretical nor actual support for the R Q as a measure of dissimilation. Since no other validation of the R Q is available at the present time, one must go further and say that there is no evidence that the R Q is a measure of dissimilation. This leaves us in

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TABLE 18 OXIDATION AND OXIDATION
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continues high for some time after intestinal absorption is complete or the injection has ceased. The total increment in the oxygen consumed (and the correspond ing extra energy expenditure) is known as the "specific dynamic action" (S.D.A.) of the foodstuff given. The magnitude of the S.D.A. differs for the different food stuffs. For carbohydrate it approximates so per cent of the calone value of the amount of sucar administered.

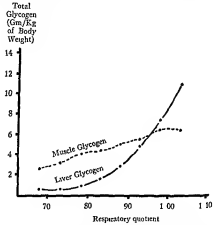


Fig. 41a -Relationship between muscle glycogen liver glycogen and R Q (Bridge [36])

Various explanations of the S D A have been advanced (22). The mechanism is undoubtedly different for each of the foodstuffs. The work of Wierzuchowski (35) is the most illuminating as regards carbohydrate. He injected glucose intra venously into dogs at rates ranging from x to 9 gm per kilogram per hour and observed the heat production, the R Q, and the sugar and lactic acid levels of blood and unne. He then correlated the S D A with his other data at all rates of glucose injection and found that there was a good proportionality between the S D A and the amount of glucose "assimilated" (the amount of glucose injected minus the

amount excreted in the utine). The glucose equivalent of the oxygen consumed was not clearly related to the S D A, nother was the fat formation, as judged by the slight use of the R Q above unity and other enteria. He therefore concluded that the S D A was related to the amount of glucose stored, which for practical purposes means the amount of glucose formed

Simultaneously with the increased oxygen consumption following earbohydrate make there is an even greater use in CO, production, so that the RQ is clevated (chap xs) Bridge (50) has pointed out a relationship between the rise in RQ and glycogen deposition similar to that found by Wierzuchowski for the SDA. Figure 414, taken from Bridge shows the correlation between the RQ and the glycogen contents of liver and muscle in a sense of riabhits at various intervals after earhohydrate administration. It will be noted that the curve relating the RQ to liver elections of the suppose of the remarkably smooth.

The work of Wierzuchowski and of Bridge suggests that the S D A or the R Q. or both, could be used as an index of glycogen formation in the intact animal or in man when the sampling of tissues is impossible or undestrable. There is a good theoretical basis for the application, quantitatively as well as qualitatively We have seen in chapter iv (see Fig. 20) that the synthesis of glycogen requires intergy which is derived from oxidative steps in the breakdown of glucose. From in vitro experiments it can be calculated that the oxidation of 1 mol of glycogen From this one might predict that the S D A of glucose would be between 8 and 17 per cent of the amount of glucose retained. The observed S D A of 10 per cent is well with in this theoretical range. It remains for future work to compare the S D A and the R Q with chemical determinations of glycogen deposition under conditions which would be feasible for chinesia use.

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PART IV

THE ROLE OF THE ENDOCRINE GLANDS IN CARBOHYDRATE METABOLISM

level of 45 mg per cent within 5 hours. One milligram of pure crystalline insulin contains 22 such units

From a historical standpoint and because of its importance as a research tool and as a therapeutic agent, insulin may be regarded as the dominant instrument in the symphony of endocrine action that results in normal carbohydrate metabolism. It should be remembered that any particular bormone is merely one of the components of the endocrine balance and that its actions depend upon the presence and simultaneous influences of the other hormones. In this sense it is difficult to deal with one hormone at a time. But, since it is even more difficult to describe the complicated actions and interactions of all the endocrine glands in a parallel fashion it does serve a useful purpose to discuss the subject as if insulin were carry ing the leitmotiv of the symphonic work while the other endocrine instruments amplified or modified the theme

THE REGULATION OF INSULIN SECRETION

In the post absorptive state and in the absence of physical emergencies or emotional crises the pancreas probably secretes small amounts of insulin into the blod continuously. This constant secretion is a prerequisite for the efficient functioning of the hepatic regulating mechanism, which is the most important factor in the maintenance of the normal blood sugar level (0, 10) (cf p 248). Hédon (11) has shown that a deficiency of insulin and a consequent rise in the blood sugar level begins immediately after removal of the pancreas. Soskin and his co-workers (12) found that it required a constant injection of insulin to maintain a constant nor mal blood sugar level in depancreatized dogs. The latter investigators further showed that no extra secretion of insulin was necessary for an adequate disposition of a sudden influx of carbohydrate (cf chap xxi, p 249). However, this does not contradict the considerable body of evidence which indicates that extra numlin it ordinarily secreted as a result of hyperglycemia following carbohydrate intake (13, 14) or as a consequence of central nervous system activity transmitted through the might vagus nervey (15, 16, 17)

It has been shown that under special experimental conditions byperglycemia may stimulate the pancreas both directly and by way of the nervous system (18 19). In the normal intact animal these mechanisms for counteracting hypergly cemia contend with other mechanisms that tend to raise the blood sugar level. For example, asphyxia and certain drugs, like metrazol, ordinarily result in hyperglycemia. In animals in which the adrenal medullae have been destroyed, these same agents cause hypoglycemia (20, 21). But when the right vagus nerve is cut in an adrenal medullectomized animal, the hyperglycemic agents produce no effect on the blood sugar level (20, 21). It is evident that vagal stimulation of extra insulin secretion acts as a restraining counterregulation in limiting the hyperglycemic effects of the adrenal medulla and the sympathetic nervous system. The

adrenal medulla and the sympathetic nervous system, on the other band, may be regarded as emergency safeguards against hypoglycemia that is too rapid or too severe to be adequately handled by other mechanisms

It is beyond the scope of this volume to discuss these emergency mechanisms in detail. It may be pointed out, however, that their peculiar status is revealed by the fact that adequate regulation of the blood sugar level (except for an increased sen sitivity to insulin) ordinarily persists even after all possible influence of the nervous system has been eliminated. This has been shown after denervation of the liver (22), denervation or gralling of the pancreas (23, 24, 25, 26, 27, 28, 29, 30), denervation or destruction of the adrenal medulla (3x, 32, 33), bilateral vagotomy (44, 34), and total sympathectomy (3x, 26).

THE KNOWN PHYSIOLOGICAL EFFECTS OF INSULIN

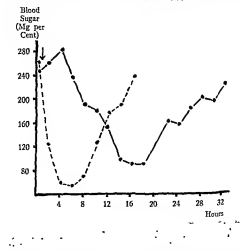
1 Hypeglycemia — Since highly punfied insulin has been available for experimental and clinical use, it has been administered to animals and burnans under the most diverse conditions. Except for differences in the magnitude of the effect obtained with a given amount of insulin (so-called "sensitivity"), a hypoglycemic effect is invariably obtained, regardless of the state of the animal. This is true for animals at any age in whatever state of nutrition, and lacking the various endocrine glands or visceral organs (37, 38, 39, 40). It is clear, therefore, that the hypoglycemic effect of insulin is a general one, which is not mediated by any particular organ or tissue. Figure 42 shows the typical curves of action of regular and of protainine insulin.

Numerous attempts have been made to determine whether the action of insulin might be on the blood itself. It has been impossible to demonstrate any change in blood in surve by the addition of insulin (41, 42). At one time it was claimed that insulin changed the blood glucose to a more reactive form (43, 44) (7 glucose), but this was never substantiated (45, 46). It is also known that insulin has no in fluence on the distribution of glucose between plasma and red blood cells (47) or on the rate of glycolysis of the blood sugar (48, 49). It seems certain, therefore, that the lowering of the blood sugar level in surve under the influence of insulin is a re sult of the more rapid withdrawal of sugar from the blood by the other insues. A decreased supply of sugar to the blood from the liver is an additive factor (50, 51, 52).

2 Glycogen deposition —Next to its hypoglycemic effect, the glycogenetic effect of insulin in skeletal muscle is its most thoroughly substantiated direct action. It is readily demonstrable in sitto on thin sheets of muscle (diaphragin or abdominal muscle of the young rat) in the Warburg apparatus (53, 54, 55). It is important to remember, however, that this action of insulin in vio is related to the existing blood sugar level from moment to moment both because of the amount of sugar available for deposition and because of the secondary counterregulations evoked.

by hypoglycemia. Thus, unless the blood sugar is maintained by the administration of sugar, the hypoglycemia resulting from insulin action will evoke a secretion of epinephin from the adrenal medulla, which, in turn, may mask the glycogenetic effect of the insulin by causing a rapid breakdown of muscle glycogen to lactic acid.

That insulin influences the deposition of liver glycogen is evident from the char acteristically low glycogen levels of the diabetic liver (56, 57) and their return to



normal with insulin treatment (58, 59). But there is a paradoxical situation as regards the effects of administered insulin in normal animals, for (with a single un explained exception [60, 61]) all normal animals invariably exhibit a decreased amount of hepatic glycogen after insulin administration (62, 63, 64). Part of this effect may be ascribed to the hypoglycemia induced secretion of epinephina and

the consequent breakdown of liver glycogen to blood sugar. But this is by no means the whole explanation for Bridge (65) has shown that insulin administered with sufficient glucose to maintain a certain blood sugar level results in a smaller deposition of hepatic glycogen than the administration of that amount of sugar alone which will reproduce the same blood sugar level. He also showed that this anom alons effect of insulin in normal animals could be obtained in the absence of the adrenal medulla.

The normal heart like skeletal muscle, deposits increased glycogen under the influence of insulin (66–67). But carefuc glycogen is apparently more dependent upon the concentration of sugar available in the blood than is the glycogen of other organs for the heart of the completely depancreatized animal may contain large amounts of it (68–69–70)—amounts which are reduced by restoring the blood sugar level to normal with muslin. The finding of Junkershoff (71) of a bigh glycogen content in the cardiac muscle of phlothizmized dogs with low blood sugar levels also suggests the possibility of the formation of cardiac glycogen in sith from non carbohydrate sources.

The glycogen content of the hrain and nervous tissues on the other hand is in finenced little if at all by either the blood sugar level or by the insulin content of the blood [27, 3]. Indeed it seems likely that the small amount of glycogen which is found in these tissues has more structural than metabolic significance, since the amount is little affected by various nutritional physiological and phar macological factors [74, 75]

- 3 Artikelogenesis—As outlined in detail in chapter x, ketogenesis in the liver is best correlated with a lack of glycogen. Accordingly insulin is antiketogenic (76 77, 78) under conditions in which it increases liver glycogen (in the diabetic organism), but it may actually be ketogenic (79 80) under conditions in which it decreases liver glycogen (in the non-diabetic organism). Insulin has no influence whatever on the rate of disposal of ketone bodies by the extrahepatic tissues (8r 82).
- 4 Change in the RQ—Whatever the significance of the RQ (chap xiv), insu in has a definite effect upon it. But the situation with respect to the difference be tween the normal and the diabetic organism and the influence of the amount of car bohydrate available is somewhat similar to that which obtains for glycogen depo a sition in the liver. Thus in the absence of insulin the diabetic organism fails to exhibit the rise in the RQ which follows the administration of sugar to the normal animal (83–84). The administration of insulin alone to the fasting diabetic organism results in an elevation of the quotient (85–86). However, insulin administration to the fasting normal organism results in variable changes of small magnitude (87–88–89), although insulin plus sugar does cause a more abrupt and more pronounced rise in the RQ than does sugar alone. Insulin has either no effect on the oxygen consumption or may actually decrease it (\$4, 55, 67, 90)

When insulin does affect the RQ, the results bear no quantitative relation to the fall in the blood sugar level According to Bridge (91), the RQ changes cor relate best with the level of hepatic glycogen (see chap nv, p 161)

5 Decrease in serum inorganic phosphate —In the absence of insulin the dia bette organism exhibits an abnormally high level of inorganic phosphate in the blood (92, 93). This is corrected by treatment with insulin (93, 94). The adminst tration of insulin to the normal animal causes a diminution of serum inorgane phosphate below the normal level (95, 96, 97). There have been variable and contradictory reports concerning supposedly parallel changes in the hexosemonophosphate content of muscle, presumally due to the entrance of the blood serum inorganic phosphate into muscle in this esterified form (98, 90). But Soskin and

TABLE 10

CHANGE IN INORGANIC PROSPHATE (P.) ANN TOTAL ACID-SOLUBLE PROS PHATE (P.) OF THE BLOOD AND IN HEXOSEMONOPHOSPHATE (HimP) OF THE MUSCLE (SOSKIN et al. [121])

(In Milliorams per Cent)

		CRANCE	CHANGE IN	
EXPERIMENTAL CONDITIONS	No No	Pe	PT	Hole*
Departerestized dogs given epi nephna (o i mg/kg subcutane- ously)	{z 2 3	-0 3 -0 1 -0 4	+30	+ 9 5 + 10 9 + 9 4
Adrenalectomized dogs given insulin (0 3 unit/kg subcutaneously)	{1 2 3	-1 9 -1 6 -1 2	†20 †30	-05 -03 +03

[&]quot; In terms of phosphate

his co-workers (42) have shown that the phosphate changes in blood and muscle are not directly related to each other and that only the fall in the blood inorgane phosphate is a direct consequence of insulin action. The confusion was due to the counterregulatory reactions, whereby excessive insulin activity evokes a secretion of epinephrin, and vice versa. When the actions of the individual hormones are isolated by excision of the counterregulating gland, the unopposed action of the administered hormone can be observed (Tables 19 and 20).

The administration of insulin to the normal intact animal is followed by both the blood and the muscle phosphate effects. In the absence of the adrenal glands, the action of insulin on the blood phosphate persists, while the hexosemonophose phate in muscle is not affected. The responsibility of reflectly secreted epinephina for the muscle phosphate changes after insulin administration also accounts for the absence of those changes in normal animals when sufficient dextrose to prevent hypoglycemia is administered with the insulin.

mal animal causes both a fall in the inorganic phosphate in the blood and a rise in the hexosemonophosphate in the muscle. But in the departreatized animal, only the muscle effect of epinephrin occurs.

The action of insulin in lowering the blood morganic phosphate is not explained by a loss of phosphate from the blood, for the total blood phosphate remains unchanged. It seems probable, therefore, that there is an esterification of the inorgan ic phosphate within the blood (42, 100), although the nature of the phosphate comnound which is formed is, as wet, unknown.

6 Decrease in serum potassium—A number of investigators have observed a lowering of the potassium content of the blood serum following the administration of insulin to normal animals (roi, 102, 103) There has been no elucidation of the

TABLE 20

CHANGE IN BLOOD INORGANIC PHOSPHATE (P.) AND IN TOTAL ACID SOLUBLE PHOSPHATE (P.) (SOSKIN et al. [42])

(In Militarius per Cont)

The maximum decrease in blood morganic phosphate (P.) obtained with glucose in any depancreatized saimal was 0 4 mg per cent. Hence no change in P. of this amount or less was connidered to be significant throughout our work

TYPE OF ASSISTAL	Gircone				Insulm			Emplyment							
	No oi	Decrease III. Av R se		se No P		P.	P. R		Av Rise No		Decrease n Pe		Av Rase		
	Dogs M	Ма	Mez	Av	Pr	Dogo	Мв	Max	Av	in Pt	Dogs	Min	Max	Á۲	In Pr
Normal Deparcreatized Adrenalectomized	5 7 3	0 5	1 2 0 4 2 8	0 8	100		0 7	2 0 2 8 2 0	1 2	10	9 14 3	0 4 0 6 1 1	17012	0 3	3 0 5 0

mechanism of this effect, except perhaps in so far as it may be related to the in creased rate of entry of sugar into tissues under the influence of the hormone Fenn has shown that potassium enters tissues in proportion to the amount of carbohydrate which is taken up (104)

7 Influence on nitrogen metabolism—In the absence of insulin the diabetic or games exerctes abnormally large amounts of nitrogen in the urine (105, 106, 107). This indicates that insulin must act to inhibit protein catabolism at some point. The in vitro work of Bach and Holmes (108) with liver slaces showed that insulin inhibits the deamination of amino acids, as judged by the decreased rate of appear acids.

similar experiments, were able to confirm this proulin effect with d alanine but not

with the naturally occurring I alanine as had Bach and Holmes. This n troven sparing effect of insulin was further demonstrated by Gaebler and others (110 III) in an indirect way They found that whereas extracts of the anterior pituitary ad ministered to normal animals resulted in nitrogen retention, the same treatment in diahetic animals caused an increased nitrogen excretion

The administration of insulin to the normal animal is followed by uncertain and contradictory results (112 113) There may be either no change or an actual in crease in nitrogen excretion. However, the amino acid level of the blood does de crease s gnificantly (114 115) Like other effects of insulin under similar circum stances this is probably due to the counterregulatory effects of other glands par ticularly the adrenal medulla Luck and his co workers (116) have shown that in adrenal demedullated animals insulin fails to lower the blood amino acids while epinephrin will do so just as it does in the normal animal. It seems reasonable to conclude therefore that the apparent influence of insulin on the amino acid level of the blood of the normal animal is actually due to the reflex secretion of epi nephrin resulting from hypoglycemia. This same sequence of events could of course also account for the increased excretion of nitrogen which sometimes fol lows the administration of insulin to normal animals for epinephrin has been shown to increase protein catabolism

However it is not at all certain that as de from secondary effects due to epi nephrin secretion insulin does not have a direct action of its own upon the blood amino acids Mirsky and his co workers (117 118) found that in eviscerated and nephrectomized does maintained by a constant injection of insulin and glucose the blood amino acids rose more slowly and injected glycine disappeared more rapidly than in similar animals maintained on sugar alone Since the absence of the liver and kidneys precludes a loss of the amino acids by deamination these experiments suggest that insulin facilitates the use of amino acids in the muscles for synthetic numoses either directly or indirectly (see chap xix p 235)

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CHAPTER XVI

THE MODE OF ACTION OF INSULIN

MORE detailed examination of the physiological effects of insulin shels some light on the manner in which insulin influences carbohydrate metab olism. It may be well to begun by directing our attention to skeleth muscle, because this tissue comprises about 50 per cent of the body weight, be cause it is a less complicated organ, in a brochemical sense, than is the liver, and because more data concerning it are available.

INSULIN AND GLYCOGENESIS IN SKELPTAL MUSCLE

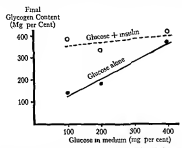
Although it is facilitated by insulin, the deposition of glycogen can occur in the complete absence of the hormone (1, 2) The fact that insulin is not essential for glycogen formation has received in wine confirmation from the work of Con and his co-workers (3, 4) They synthesized glycogen from glucose in the test tube in the presence of the necessary enzymes but without insulin Indeed, they were unable to demonstrate any effect when insulin was added to their system (5 6) In the living animal, Dambrosi (7) and Lukens (8) have shown that the absence of insulin does not even limit the extent to which glycogen is restored after its depletion by exercise. It is the rate of restoration of glycogen which is deficient, for, whereas in the normal animal it took x hour to restore the pre-existing glycogen level, the muscle glycogen of the completely deparcreatized animal was restored just as fully in 4 hours Insulin, therefore, exerts its influence on the rate of glycogen formation

The major factor, other than insulin which determines the rate of glycogen synthesis is the concentration of sugar present. This is, of course, in accord with the general nature of all enzyme reactions. Core it all (s) have shown that the amount of glycogen deposited in the liver of a given experimental animal depends upon the height at which the blood sugar level is maintained rather than upon the total amount of sugar given It has been possible in our own laboratory (is) to demonstrate this relationship for muscle even more clearly on rat disphragin is vitro by the Warburg technic. Figure 43 shows the increasing amounts of glycogen deposited at increasing sugar concentrations, with or without added insulin. It will be noted that at the highest concentrations of sugar the insulin exerted no significant effect over and above the effect of sugar concentration. This relationship of insulin action to sugar concentration is consistent with other actions, which are to

be discussed later. In other words, insulin enables the tissues to do at low or physiological sugar concentrations that for which they would otherwise require very high sugar concentrations.

INSULIN AND THE UTILIZATION OF SUGAR BY SKELETAL MUSCLE

One of the most firmly intrenched notions about insulin in the metabolic literature is that it increases the dissimilation of carbohydrate. This is without basis in fact, for, as pointed out in chapter xiv, no over all measure of dissimilation in the living organism is yet available. The supposed proof for the mistaken asser



Fro 43.—Influence of sugar concentration on deposition of glycogen in rat diaphragm in vitro with and without insul n (Hechter et al. [10])

tions is based upon calculations of so-called oxidation" from the R Q (see chap xi) and the estimation of utilization from carbohydrate balance experiments

Wierzuchowski (11) used R Q measurements to calculate the amounts of sugar oxidized before and after the administration of insulin in two normal unanes thetized dogs receiving constant intravenous neuterions of glucose (Table 21). Ac cording to these calculations one of the animals "oxidized 21 for cent of the as similated sugar before insulin administration and a7 3 per cent after insulin. But the other animal "oxidized" 19 1 per cent before insulin and 19 o per cent after insulin. The results from the two dogs were averaged, and the conclusion arrived at was that insulin bad increased the oxidation of assumilated glucose from 20 3 per cent to 23 a per cent!

Best, Dale, Hoet, and Marks (14) measured ovygen consumption and made carbohydrate balance studies on the same eviscerated spinal cats. In the absence of the liver they found that an increased amount of sugar disappeared from the blood following insulin administration and that the sugar which disappeared was equal to the sum of the glycogen deposited in the muscles and the glucose equivalent of the oxygen consumed In accordance with the state of knowledge at that time, Best et al. (15) concluded that the effects of insulin in excess present ain tensification of its physiological effects, including the acceleration of the combustion of carbohydrate. Hence their work has since been quoted as proof that insulin increases the dissimilation of carbohydrate.

A re examination of their original data shows that this conclusion was not war ranted. Table 24 summarizes the periment figures from the experiments which they themselves selected as being most free from technical criticism. The right

TABLE, 24

INFLUENCE OF INSULIN ON GLUCOSE OXIDATION OF EVISCERATED

SPINAL CATS (BEST & al. full)

	RECALCULA TION				
Exper ment No	Insul a (Units)	We ght of Cat (Rg)	Duration of Experiment (Min)	Glucose Ox d sed (Mg)	Glucose Oz d sed (Mg/Eg/Ht
5A 5B 0	0 20 30 25	3 2 3 2 2 6 2 8	50 150 210 250	1 045 2 970 2 595 3 079	392 371 285 264

hand column is our own recalculation of the amounts of sugar "oxidized' in milli grams per kilogram per hour, inserted in order to make these values comparable It may be seen that animal No 5 "oxidized" less sugar after insulin than before Animals Nos 6 and 7, for which no pre insulin periods are given, "oxidized less sugar after insulin than animal No 5 "oxidized' without insulin It is clear that this work offers no support for the contention that insulin increases the rate either of utilization or of the so-called "oxidation of carbohydrate".

The more recent work of Soskin and his co workers (16, 17) has confirmed the fact that insulin does not increase the utilization of carbohydrate in the organism as a whole, while at the same time giving some insight into the reasons for the previous confusion The form of the experiments was a chemical balance study in liverless dogs, as described in chapter xiv, where the relation of carbohydrate utilization to blood sugar level was discussed Experiments similar to those which were done on the normal animals, were repeated on completely depancreatized dogs which had been deprived of food and insulin for 3 days Figure 44 summarizes

the results and compares them to those obtained in normal animals. It may be seen that dextrose utilization in the depanceratized dog is qualitatively similar to that in the normal dog. In both types of animal the rate of utilization depends upon the height of the blood sugar level. Within a widerange of blood sugar values the diabetic dog utilizes less sugar than does the normal dog at any particular elevel. But, above certain high levels this difference disappears and both types of animal use the same amounts of carbohydrate at the same blood sugar.

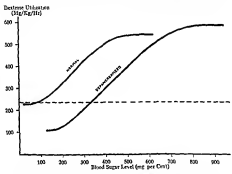


Fig. 44 — Relationship between blood sugar level and dextrose utilization in normal and in depan creatized dogs (Soskin and Levine [16])

levels. When, however, one compares the rate of utilization of the normal animal at its usual normal blood sugar-level with the rate of utilization of the diabetic animal at the hyperglycemic levels which it ordinarily maintains, it is apparent that the diabetic animal habitually uses as much or more sugar than the normal animal.

It is also clear that, when one administers insulin to a diabetic animal, two mutually counterbalancing effects are obtained there is a potential increase of the amount of carbohy drate that can be utilized at the pre-existing blood sugar level, but there is also a coincident reduction in the level. The net result is no change in the tate of utilization. In view of these results, insulin cannot be regarded as essential to the utilization of destrose or even as a determining factor, so far as the net result is concerned. It apparently plays the part of a catalyst or activator in a process which can proceed at a slower rate in its absence. More specifically, it per mits rates of carbohydrate utilization at low blood sugar levels which in its absence would require abnormally high blood sugar levels.

The question then arose as to whether the amounts of insulin available in the normal animal were such as to result in maximal rates of utilization at any given blood sugar level. To answer this question, carbohydrate balance experiments were performed on existented normal animals maintained at particular blood sugar levels despite the constant administration of large amounts of insulin (17)

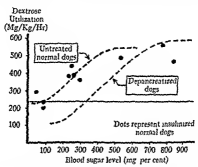


Fig. 45 "Influence of adm n stered insulin on sugar utilization in normal dogs. (Soskin and Le tine [27].)

the enr

normal animal is already optimal as regards the utilization of sugar, so that audit tional insulin causes no change. But this is not the case as regards the storage of muscle glycogen, which is increased as a result of additional insulin.

Considering the fact that the lack of insulin causes a diminition in both whilst tion and storage of carbohy drate by the peripheral tissues at any given blood sugar fevel, it seems probable that insulin acts by promoting the conversion of gluces into some intermediate substance which is necessary for both processes. It may be supposed that the rate of formation of the intermediate substance depends upon

the concentration of the blood sugar and upon the amount of insulin present In the untreated depanceatized animal the increased concentration of blood sugar can by itself lead to the formation of sufficient intermediate substance to support the normal rate of catabolism. However, there is little or no excess of the intermediate substance available for synthesis to glycogen. The administration of in sulin to the depanceatized animal increases the amount of intermediate substance formed at any blood sugar level. The animal now resembles the normal in having available sufficient intermediate substance to maintain the normal or maximal rate of catabolism even at normal blood sugar levels. There is now also available additional intermediate substance for synthetic purposes. In the normal animal, in

TABLE 25
INFLUENCE OF INSULIN, IN RELATION TO THE BLOOD-SUGAR
LEVEL, ON THE RATE OF ENTRY OF GLUCOSE INTO THE
PERIPHERAL TISSUES OF LOVELISES AVAILAGE

Bases-Steam	Minuscanus Traces			
Martaines (Me/100 Cc)	Departures Sland	Normal	Normal+ Added in- sulis	kimisa
45 80 160 230 525 020 750	79 50 355 415 471	15 104 115 161 400	340 577 578 491	Soskia and Levine (16, 17)
200 240 325		124 150 200	4% 1 %	Best et al (14 15)

which sufficient intermediate substance is already present to allow the catabolic reactions to proceed at their maximal rate, additional intermediate substance resulting from insulin administration is reflected only in increased glycogen synthesis

If the action of insulin in the tissues is to promote the conversion of glucose into some intermediate substance which is necessary for both utilization and glycogenesis, a consistent effect of the hormone should be an increased rate of entry of sugar into the tissues, regardless of the fate of the sugar thereafter. Ample data to show that this is the case were furnished by the carbohy drate balance experiments, in which sugar was constantly injected in order to maintain constant blood-sugar levels (16, 17). Table 25 summanizes these data, as well as the results of comparable experiments of Best et al. (14, 18, 18).

INSULIN AND THE DISSIMILATION OF CARBOHYDRATE BY SKELETAL MUSCLE IN DITO

Since the advent of the Warburg technic, there have been a number of aitempts to demonstrate the action of insulin in witho. These attempts have been successful as regards the deposition of glycogen in isolated muscle (p. 169) but have been uniformly unsuccessful in showing any influence of insulin on so called 'oxidation or dissimilation of carbohydrate in mammahan muscle (18-19-20). As in the whole animal, insulin causes either no change or an actual decrease in oxygen consumption, and there is no correlation between the oxygen consumed and the sugar which disappears (Table 18, p. 157).

In contradistinction to the results obtained in mammalian muscle, Krebs and Eggleton and others (21, 22) were able to demonstrate an increased oxygen con sumption under the influence of insulin in minced pigeon breast muscle. These experiments were performed in the presence of glucose as the substrate and with the addition of citric acid as a catalytic agent. The high R.Q. obtained under these circumstances led to the conclusion that the increased oxygen consumption resulting from the addition of insulin signified a stimulation of carbohydrate "oxidation" by the bormone. Using the same tissue and pyrivate as the substrate, Rice and Evans. (33) demonstrated an increased oxygen consumption with a coincidentally increased disappearance of pyrivate under the influence of insulin Apparently, an insulin effect on some oxidative process is obtainable in pigeon breast muscle.

The work on the muscle of birds tends to confuse the picture of insulin function rather than to clarify it for it must be pointed out that, of all the experimental auxiliars pigeons are about the least suitable from which to draw conclusions of general significance. It takes relatively enormous doses of insulin in the intact bird to produce even a small fall in the blood-sugar level. On the other hand, the removal

of the fate of pyruvate in manimalian muscle According to Flock and holination (a6), administered pyruvate is disposed of just as rapidly by the completely depancreatized dog as by the normal dog, while Bueding and Himwich (27) have shown that the administration of insulin with glucose actually results in a greater rise of pyruvic acid in the blood than does the injection of the carbohydrate alone. In view of these facts, it seems necessary to reserve the work on pigeon heast muscle for future interpretation, for it is impossible to correlate or reconcile the results in birds with the much larger hody of information obtained from manimals

THE INFLUENCE OF INSULIN ON THE LIVER

Evidence as to the mode of action of insulin in the liver is less abundant than the evidence for muscle, chiefly because hepatic tissue appears to be so sensitive to environmental factors that relatively few successful in vitro or perfusion expen

ments have been reported. When studying the intact living organism, the results are difficult to interpret because of the many regulatory and counterregulatory in fluences to which the liver is subject. Nevertheless, there are sufficient data to show that the actions of insulin on the liver are correlated with the blood sugar level, as they are in muscle.

Issekutz and Szende (28) were the first to demonstrate that insulin inhibits hepatic glycogenolysis. They showed that livers removed from frogs which bad previously received insulin produced less sugar than did the livers of untreated frogs. Similar, though less well-controlled, results were obtained by Cori (29), Molitor and Pollak (30), and Sahyun (31) by different methods. On the other hand, Lundsgaard et al. (32, 33) were unable to show that insulin had any action on glycogen hreakdown or deposition in the perfused livers of cats and dogs

More recently, however, Soskin and his co-workers (34, 35) were able to demonstrate an inhibitory effect of added glucose on the rate of appearance of free sugar in mined dog liver in wire. This offered the opportunity for the testing of the action of insulin on hepatic glycogenolysis under simplified conditions. A lobe of the here was removed from normal dogs anesthetized with nembutal Insulin was then administered to the animals, and 30-45 minutes later the remainder of the liver was removed. In the liver samples removed after insulin administration, there was a significantly lower rate of appearance of free sugar than in the samples removed before insulin was given. When glucose was added in vito to hoth sets of liver samples, the rate of glycogenolysis was inhibited to a greater extent and hy smaller amounts of added glucose in the "insulinized" samples than in the control samples (Table 46). It was apparent that insulin inhibited glycogenolysis in the liver and reinforced the inhibitory effect of added dexires.

The antiketogenic and nitrogen sparing effects of carbohydrate are ordinarily considered as requiring the presence of insulin, since they are difficult to elect in its absence. But the hormone is not essential, as has been shown by Soskin (26) In fact, this work demonstrated that every criterion of carbohydrate utilization which is exhibited by the normal annial can also be obtained without insulin in the completely depanceratized animal under the appropriate experimental conditions (see p. 107). More recently, Mirsky and his co-workers (37, 38) have shown that the antiketogenic and nitrogen sparing effects of carbohydrate can be obtained in a cutiley diabetic animals without insulin if the blood sugar is raised to a sufficiently high level. Hence, the mode of action of insulin with respect to the foregoing he-

PROSPHORYLATION, THE COMMON PACTOR IN INSULIN ACTION

It is a reasonable a priori assumption that the various physiological effects of insulin do not represent different and unrelated functions of the bormone. It is

more likely that they are indirect consequences of a single catalytic influence on some basic enzyme system. From the functional standpoint the fundamental action of insulin may be considered as being the increased rate of entry of glucos (dextrose) from the blood and extracellular fluids into the tissue cells of the body This may not apply to organs like the brain and kidney; but it does apply at feast to the skeletal muscles and liver, which compose the overwhelming bulk of the metabolically active tissues of the body In biochemical terms, the increased transit of sugar into the tissues may be described as the facilitation, by insulin, of a

TABLE 26
INHIBITION OF GLYCOGENOLYSIS IN LIVER BREI BY DEATROSE AND BY
DEXTROSE PLUS INSULIN (TABBENHAUS & a) [1:t])

EXPT No Time Cassort parts	_	Total	ANOUNT		SUGER®	Percentage of Incident		
	DERTROIR Annes	Without fasuls	Wath Insulin	Veith Dextrose Alone	With Dextrose +Insulin			
1	60 60 60	2,778	04 186	141 299 319 267	165 145	0	45 52	
It	0 0 0 0 0 0 0 0	3,313	0 100 203 418 835	254 1,997 2,027 1,565 1,229 1,260	1,180 1,073 305 661	0 22 39 37	42 46 60 61	
ш	60 60 60 60 60	4,184	0 100 200 450 900	105 1,317 1,196 1,080 1,019 446	217 2,269 2,066 892 690 222	9 18 23 65	19 32 48 83	

^{*} Values are in miligrams per 200 gm of liver calculated as for glucose

basic phosphorylation which introduces carbohydrate into the metabolic processes of the cell Regarded from the physical aspect, it may be said that, by increasing the rate of phosphorylation of glucose within the cell, insulin causes a steeper gradient of free sugar across the cell membrane and thus increases its rate of diffusion into the cell

As outlined in chapter iii, the present state of knowledge of the intermediate steps in carbohydrate metabohism indicates that the intermediate substance, the formation of which is facilitated by insulin, is one of the phosphorylated hexoss It will be recalled that the phosphorylation of sugar in the cell is accomplished by a substance possessing high-energy phosphate bonds, namely, adenosine triplos-

phate (ATP) The original energy necessary for the production of ATP from adenylic acid must eventually come from such oxidative reactions as may be coupled with the esterification of inorganic phosphate. It must therefore be assumed that insulin acts at an as yet unknown locus in this cycle of events (39-40). This is consistent with the demonstrated effect of insulin in esterifying inorganic phosphate in the blood (p-172). It is also supported by the recent work of Sacks (41) with radioactive phosphorus in which he showed that insulin increased the rate of tumover of the phosphate in ATP.

This hypothesis is compatible with the observed relationship between insulin action and sugar concentration. It is to be expected that the rate of the hasic phosphorylation, like that of any other enzymatic reaction would be influenced by the concentration of the substrate. Like any other catalyst insulin could be regarded as increasing the rate of the reaction for any given concentration of the substrate if the substrate concentration were high enough no additional effect of the catalyst could be demonstrated. At low concentrations the action of the catalyst could be described as making possible rates of reaction which in the absence of the catalyst would require very high concentrations of substrate.

From the foregoing point of view the various physiological effects of insulin which have been described as separate phenomena emerge as merely different parts of the same chain of events. The fall in the blood sugar level is a direct reflection of the influence of insulin on the basic phosphorylation in so far as it causes a greater rate of removal of sugar from the blood. The association of notassium with the hexose phosphates in muscle also accounts for the withdrawal of blood potassium. The accelerated metabolic processes made possible by the in creased rate of the first step in the series results in a greater disposal of the substrate both for synthetic and catabolic purposes (glycogen deposition and R O change) The increased availability of the substrate to the enzymatic machinery of the cell allows carbohydrate to become predominant over protein and fat in the competition for the oxidative systems. Hence the cataholism of protein and fat is inhibited (antiketogenesis and mitrogen sparing action) The latter effects are nat urally prominent in the liver which is primarily concerned with the interconver sion of foodstuffs while the former effects are more characteristic of the skeletal muscles and other effector organs which derive their energy chiefly from carbohy drate and ketoacids

INSULIN AND THE ENZYMATIC MACHINERY OF CARBOHYDRATE METABOLISM

The dominant role of ATF in tissue phosphory lations was described in chapter in This high-energy phosphate compound is formed from adenylic acid and mor game phosphate and the potential energy which it represents and which must be forthcoming for its continuous formation is presumably derived from oxidative steps in the dissimilation of carbohydrate (Fig. 46) Using radioactive phosphorus

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TABLE 26

Inhibition of Glycogenolisis in Liver Brei by Dextrose and by Dextrose are India to Claderolisis and Lot.

Exer ho	{ _	TOTAL	Awoput		SUGAL*		PERCENTAGE OF	
	Ther.	CARBORN	DEXTROSE ADDED®	Without Issal v		W th Dextrose Alone	N b Dextross +Iosulu	
1	•	2 778	•	141				
	60 60 60	1		299				
	60	1	186	310	165	0	45 52	
	60	2 897	186	267	145	11	52	
11	۰	3 313		254				
	60		101	1 997				
	8888	J	100	2 027	I 180		42 46 60 67	
	60	ſ	208 (I 565	1 073	22	45	
	60		418	1 229	805	39 37	60	
-	60	3 410	100 208 418 836	1 250	662	37	67	
111		4 184		205	217	1		
	60		•	T 317	1 259	1		
- 1	60		100	1 100	1 066	9 1	19	
	60	. !	200	1 080	89 z	18	32	
	6a 6a 6o		450	1 010	6ga	9 18 23 66	19 32 48 83	
ſ	60	4 000	900	446	272	66	83	

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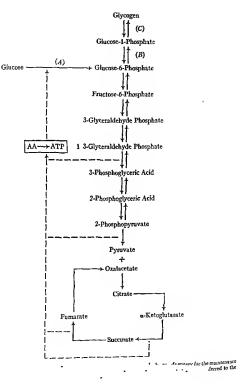
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as a tracer, Sacks (41) has demonstrated a more rapid turnover of ATP in the skeletal muscle of intact animals when glucose and insulin were administered than when glucose alone was given. Since the rate of phosphorylation of glucose depends upon the rate of turnover of ATP, it is obvious that insulin might act on any of the condutive reactions that supply the energy for the rephosphorylation of adenylic acid. But the fact that insulin does not increase oxygen consumption, either in wio (42) or in wito (Table 27), makes this simple explanation untenable. This anomalous situation might he revolved by supposing that, without actually increasing the rate of oxidative reactions, insulin increased their efficiency as regards phosphorylation so that more moles of inorganic phosphate were esterified per mole of oxygen consumed (40). This is not unreasogable, in view of the fact that different investigators have reported various rates of phosphate esterifica

TABLE 27

LACK OF EFFECT OF INSULIN ON THE OXYGEN CONSUMPTION
OF MAMMALIAN MUSCLE in pite

Cond tion of Animal	Type of Taue	Glucose in Medium (Mg per Cent)	Roula	Ço,	Reference
Normal	Abdominal muscle	400 0 0	++00	3 °) 3 °) 3 °)	Levine et al (39)
Normal	Diaphragm	200 200 200	000+	4 7 4 7 5 3 4 7	Gemmili (20)

tion to oxygen consumption according to the experimental conditions which they employed

Considering the fact that insulin usually raises the R Q without affecting the overconsumption, one might suppose that insulin acted on some as yet un known non orudative decarboxylation. But this could hardly be a direct or essential part of insulin action, in view of certain of Germaill's results (Table 28). It may be seen that he was able to demonstrate a very agmilicant action of insulin as regards glycogen deposition with no appreciable influence on the R Q.

Of course insulin might act higher up in the scheme of dissimilation and be concerned either with the enzyme acting directly on glucose (Fig. 46 step A) or with the systems between glucose 6-phosphate and gly cogen (Fig. 46 steps B and C). It has been possible to test the latter systems with purified enzymes in ritro, and the results have been negative as regards any effect of insulin (3, 6). It has like wise been shown that in the absence of added glucose insulin has no effect upon the rate of glycogen breakdown in mammalian muscle in vitro (Fig. 47). Unfortunately, it has thus far been impossible to obtain an extract of skeletal muscle which will phosphorylate glucose in vitro. An enzyme obtained from yeast and known as "hexokinase" will do so, but it is not influenced by misulin. However, hexokinase need not be similar to the enzyme system responsible for glucose phosphorylation in mammalian muscle, for, while hexokinase will phosphorylate fructose even

TABLE 28

Influence of Insulin in Increasing the Deposition of Glycogen in Rat Diaphragm in with without Affecting the R O Significantly

Glucone in Med um (Mg per Cent)	Insul a	Q0,	Total Carbo- hydrate Change n Tusue (Mg/100 Mg)	R Q Reference	
200 200 500 0	+0+00	4 8 4 9 4 6 5 2 4 7	-0 09 +0 37 +0 82 +0 56 +1 18	0 73 0 86 0 01 0 86 0 86	Gemmill (19 20)

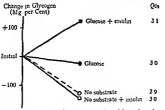


Fig. 47 —Lack of influence of insulm on glycogen content of rat abdominal muscle in a tro in the absence of glucose (Levine et al. [39])

more readily than it does glucose, mammalian skeletal muscle in vitro will deposit glycogen only from glucose and not from fructose, mannose, or galactose (Fig 48) Until the glucose-phosphorylating enzyme of muscle is isolated, it will be in possible to decide whether or not insulin may set at this point

Another difficulty is our present uncertainty as to the correctness of some of the details in our conception of carbohydrate dissimilation as outlined in Figure 46 For example, in the same experiments in which Sacks (41) showed that insulin in

creased the rate of turnover of ATP, he was unable to find any corresponding in crease in the rate of turnover of glucose 6-phosphate. This may mean that, in schetal muscle, glucose is phosphorylated to glucose i phosphate rather than to sacietal musice, guiose is phosphioryrated to gondoe a phosphate rather than to glucose 6 phosphate. If this were so, muscle would differ from brain, liver, and kidney, the extracts of which have been shown to phosphorylate glucose to glu

Data of more positive significance indicate that the point of action of insulin is probably above pyruvate. In the presence of sodium fluoride, which inhibits so phousely above pyravate in the presence of socium muonoe, which inmuts glycolysis at the phosphoglycene acid stage, the addition of insulin to muscle cose 6-phosphate gycorysis at the phosphogycenic acid stage, the addition of minim to musue in ritro still leads to a greater estenfication of morganic phosphate (Fig. 49). A in three sum leads to a greater externication or morganic phosphate (Fig. 49). A similar significance may be attached to the recent work of Himwich et al. (27) on annuar agrance mey be assessed to the secent was a sampled con (41) on department and in the blood after department dogs. They found a greater rise of pyruvic acid in the blood after



Fig. 45—Lack of format on of gl) cogen from fructone as rat abdominal muscle in a 180 with or with a handle of the control of

the administration of glucose plus insulin than resulted from the giving of the out insulin (Levine et al 1301)

The work of Bach and Holmes (43) on later shees in vilno in which they demon THE WORK OF LIBERT BIRD TROUBES 443) OF ENERGY SUCCESS IN 1419 IN WHICH THEY GETTION STREET, THE STREET BIRD TO A STREET BIRD TO STREET BIRD same amount of glucose alone satated that insume innuited ocanimation suggests a rocus of action of insume en-tirely outside of carbohydrate metabolism. Taken at its face value, this work urely ourside or carnonyurate increasional. Assett at 113 1800 value, (into work could mean either that meulin has more than one fundamental action or that it neces protein mercanonism unectry and carbonyurate meranonism only matricity library, it seems more blely that the reverse of the latter is the case. Insulin may produce this effect not by any direct action on the amino acid ondase but by in creasing the rate of entry of carbohy drate into the metabobic cycle

asing the rate of entry of earoon) drate into the metabolic Cycle. As regards a possible direct influence of insulin on fat metabolism, which might As regards a possible direct inducate of distant out at the troublem, which might be predicated on the basis of its notable antikelogenic action in the intact animal, or predicated on the mass of its indiante annoccongenic action in the intact animal, there is no pertinent in vitro work available. The enzyme systems concerned with fat metabolism are almost completely unknown

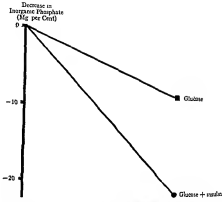


Fig. 49—Influence of insulin on the decrease in inorganic phosphate (esterification) in rat abdominal muscle in vitro (Levine et al. [39])

TABLE 20

LACK OF SIGNIFICANT DUPPERENCE BETWEEN NOWALL AND DIABETIC MUSCLE in 18th For the respiratory expeniments intact abdominal muscle of young rate (60-80 gm) was used The phosphate partitions were determined on the gastrocinemio of the stance animals P == morganic phosphate, P == two thirds of adenosine polyphosphate, P Creat == creating phosphate, P Total = total and soluble phosphate.

CONDITION	No or Anneas	BLOOD SUGAR (Ma	Qo.	Lacric Acts Proservor (Ma Pra 100 On R Q Ge era fit) Figural Partition (Mo Pre Cert)						XM
		CENT)			In O.	In Na	P.	P,	P Crest	P Total
Normal Diabetic (alloxan)	10	123 393	3 8 3 2	o 81 o 78	78 83	355 278	17 72	32 34	\$5 57	139 143

It is evident that while we are perhaps closer to the solution of insulin action than we are to the action of any other hormone, the problem is far from solved It may be that the failure to arrive at the ultimate solution depends upon the fact that very little in vilro work has been done with the tissues of diabetic animals. We have seen that in the normal intact animal an optimal amount of insulin, as regards glucose utilization, is present, so that the administration of additional in sulm is without influence in this respect. It does increase glycogen deposition in the muscles and this effect can also be demonstrated in isolated normal muscle an vitro. Further an vitro investigations using diabetic instead of normal tissues might be fruitful as regards other influences of insulin Certainly, not much prog ress was possible in the search for the points of action of the various components of the vitamin B complex on the enzymatic machinery of metabolism until the tissues of animals deprived of specific vitamins became available. However, the authors must confess that their own in piles studies with diabetic mammalian muscle have not been very enlightening thus far Table 20 indicates a number of re spects in which the diabetic muscle does not appear to differ from normal muscle

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CHAPTER XVII

THE ADRENAL CORTEX

HE essential nature of the adrenal glands to the well being of man was first indicated in Addison's original description (1) of the disease which has since been called by his name. The influence of the gland on carbohy drate metabolism was for a long time ascribed to the secretion of the adrenal medulla. In 1909 Porges (3, a) reported the occurrence of bypoglycemia in Addison's disease, which was by that time recognized as primarily affecting the adrenal cortex. He also demonstrated the occurrence of low carboby drate levels in bilater ally adrenalectomized dogs. Despite subsequent substantiation of these findings, little advance in knowledge as to the carbohydrate functions of the adrenal cortex was made until the early work of Britton and his co-workers (4, c, 6).

Stewart and Rogoff (7) had previously made adrenal cortical extracts capable of maintaining the life of adrenalectomized animals. Swingle and Pfiffiner (8, 9) die wied a new method for extraction but were particularly struck by the influence of their extract on salt and water metabolism. Britton and his co workers (5, 10, 11), on the other band, emphasized the importance of hypoglycomia and low glycogen levels as factors leading to the death of their adrenalectomized animals. While they also observed certain effects on the sodium and potassium levels of the blood, they insisted that the prepotent influence of their extracts was exerted on carbo bydrate metabolism.

The controversial nature of the subject gradually abated as it became apparent that both the mineral and carbohydrate effects were sahent features of adrenal ectomy, that they could be obtained with adrenal cortical extracts, and that they were not completely independent of each other. The use of depancreatized and of hypophysectomized animals facilitated the establishment of the carbohydrated functions of the adrenal cortex, and, finally, the potent steroids, separated from the extracts by Reichstein (12) and by Kendall (13), have made possible the accumulation of data on each aspect of adrenal function, uncomplicated by the other

THE STEROIDS OF THE ADRENAL CORTEX

The attempts at isolation of the adrenal cortical hormone have made it sufficient by evident that, whatever its natural structure, extracts of the gland may not be regarded as containing a single active substance. Hartman et al. (14) have reported that they find two factors in adrenal cortical extracts which potentiate each other but which have largely separate actions. One maintains the sodium levels of the tissues but is relatively ineffective in maintaining appetite and normal behavior and in preserving life in adrenalectomized cats. The other factor ("cortin") is very potent in preserving life, appetite, weight, and normal behavior even while the serum sodium remains low. In the light of other work, however, the views of Hart man et al. would seem to represent an oversimplification of the problem and to

17 HYDROXYDEHYDROCORTICOSTERONE

Fig to -Representative steroids of the adrenal cortex

minimize the importance of the sodium and potassium balance for the well being of the living organism

The isolation and identification of a number of steroids (13-15, 16) from the adrenal cortex and the study of their physiological properties and those of the amorphous fractions have revealed that the various compounds or fractions have certain activities in common. Figure 50 shows the formulas of some representative cortical steroids. However, a particular compound or fraction may exhibit one activity to the highest degree and be relatively impotent in other respects. In the ab

sence of more precise knowledge of that vital fuoction, the failure of which is the most urgent cause of death in untreated adrenalectomized animals, it is convenient to compare the various cortical steroids and fractions in regard to the following effects on such animals: (a) the maintenance of life, (b) the restoration of normal activohydrate levels in all tissues, and (c) the restoration of normal sodium and potassium balance and exerction. To these effects may be added the restoration of the ability of the muscles to continue to perform work in response to prolonged stimulation, according to the test developed by Togle (r₂). But since the activities of substances in this respect run parallel with their carbohydrate effects, these two actions may be considered together.

Kendall's amorphous fraction (cortin) and his desoxy B compound seem to be the most notent for maintaining life (18, 10) The carbohydrate levels are best re stored by corticosterone and its derivatives which have an oxygen or hydroxyl group on C11 (19, 20) In this respect, cortio has some effect, but desoxycorticosterone has very little (21) The relative potencies of the substances acting on carbohydrate levels maintain a similar relationship when these materials are tested on muscular work performance (10, 22) Some of the earlier work with synthetic desoxycorticosterone acetate, while showing its powerful influence on the sodium and notassium halance, had revealed no action on carbohydrate metabolism (22. 24) This is apparently a matter of dosage, for Harrison and Harrison (25) have reported that 1 25 mg daily of the substance would maintain life and a normal mineral halance in adrenalectomized rats but that it required 2 5 mg daily to maintain a normal blood sugar level Similar evidence is available in the work of Britton and Corey (26), Ingle (27), Wells (28, 20), and Long, Katzin, and Fry (21). although these authors differ from Harrison and Harrison and from each other as to the comparative potency of desovycorticosterone on carbohydrate metabolism

Table 30, which modifies and amphifies one of Ingle's (19), summarizes the relative quantitative effects of salt and of steroids, which have been shown to substitute for the functional activity of the adrenal cortex in one respect or another

DEFICIENCES RELIEVED BY SALT TREATMENT

In spite of the qualitative difference in the prepotent activity of the vanous substances which may be separated from adrenal cortical extracts, it is impossible to discuss the materials concerned with the metabolism of the foodstudits without also considering those which primarily affect the material balance. This is because the absence of the latter in adrenalectomized animals disturbs the normal environment of all cells and thus produces certain secondary disturbances in metabolism. The secondary effects are most readily distinguished from the primary metabolic effects of adrenalectomy by a consideration of those disturbances which are allevit atted by combating the mineral imbalance with a high sodium and low potassium

intake. The symptoms of adrenal cortical insufficiency which are relieved by salt treatment are as follows

- 1 Decrease in the sodium content and increase in the polassium of the blood serum— This is accompanied by an increased excretion of sodium in the urine and a decreased excretion of potasium (ao, 3;1 The changes in excretion are known to be due to a specific effect upon the kidney tubules (2;) The changes in the blood levels are due partly to disturbed kidney function and partly to a similar derange ment of electrolyte balance in the other tissues of the body (12, 3).
- a Dehydration and hemoconcentration —These are secondary to the loss of H₂O involved in the excessive excretion of NaCl They are partly responsible for the

TABLE 30

DEGREE OF RESTORATION TO NORMAL OF THE EFFECTS OF ADRENALECTOMY
BY VARIOUS MODES OF SUBSTITUTION THERAPY

(++++ = Complete Restriction)							
	Degree of Restoration by-						
CONDITION	NaCi	Desoxycor- 1 costerone	C. Sterolds				
Low blood NaCl High blood potateum Survival on food Low basal metabolic rate High blood urea Low caseboydrate absorpt on Survival on fasting Low caseboydrate absorpt on Survival on fasting Low resistance to feet glecose Low resistance to stress Low integer extretion on fasting Work performance frought measurivity Low careboydrate levels on fasting Reciscum of diabetic hyperglycemas and glycosults	++++++++++++++++++++++++++++++++++++++	***** **** **** *** *** *** *** ***	**************************************				

- rise in blood urea, although the disturbance in kidney function also contributes to this effect (34-35)
- 3 Acidosis This is due to the retention of acid metabolites and amons, which are ordinarily neutralized and excreted by the kidneys. The failure in excretion is due in part to the circulatory failure and in part to the specific kidney disturbance. A feature of the latter is an inability to produce NH₃ for the regulation of the acid base balance.
- A Impairment of carbohydrate absorption by the gastro intestinal tract and of the glycogen deposition from ingested carbohydrates—These effects may be related to the movement of potassium out of all tissue cells. Fenn showed that the passage of sugar into the cell was accompanied by a movement of potassium in the same direction (30).

- 5 Decreased metabolic rate -This has been demonstrated for the isolated tis sues of adrenalectomized animals in vito (37, 38). In the hing animal it may also sues of adirenal extensions annuals 18 vitro (\$7,30). In the name aminor is may also depend upon the reduced blood chloride level, which interferes with the dissocial uspens upon the structure most thousand ever, which interferes with the cussodar flon of oxygen from oxyhemoglohm, decreasing the supply of oxygen to the tis sues (39, 40)
- 6 Anorexia and the consequent lack of gain in weight and cessation of growth-No explanation for the loss of appetite is available
- 7 Rapid deterioration and death of the animal This is prohably a result of the 7. Adjust accentration and acoust of the unimate — Little is providing a result of the cumulative effects of dehy dration and hemoconcentration leading to a shocklike condition, plus the toric action of high potassium levels and the hypoglycemic ef fects of fasting owing to the anorexia

The beneficial effects of salt on the above symptoms are striking and very read And concentrate of soil on the above symptoms are stituing and very read ily demonstrated. The diminished rate of glucose absorption is completely reay utanusatated and diministration of NaCl in the dimking water (41, 43). The some to normal my the aumandmenton of tract in the unmaning water (41,447) and same holds true for fat absorption (43). Similarly treated advanalectomized rate same nous true nor rat, accomption (44) summary treated automatecromized rats can deposit glycogen from glucose nearly as well as normal rats (42, 44) and may gain can ucposit grycogen itom grucose nearly so weiliao normal fato (42 44) and may gain weight in normal fashion (45). But while salt treatment enables adrenalectomized animals to survive indefinitely under favorable conditions. It does not restore them completely to normal. They are still sensitive to stresses and strains of all kinds (19 44) Nor is this sensitivity completely abolished by treatment with the ster (19 44) NOT 13 this sensitivity completely accurate by treatment while the size olds that are active as regards carbohydrate metabolism (10 34) It is upon this ous that are active as regards caroony mate metacous in (19-34). It is upon this evidence that the possibility of the existence of a separate life maintaining" prin

The observations of the normal absorption of catbohydrate and fat in salt and observations of the normal absorption of cathonyurate and lat in said traded adrenalectomized animals (44) are directly opposed to the theories of Ver are uncountered animals (44) are uncountered opposed to the theories of ver are This author, starting with his observation that the intestinal absorption of the zar ams autor, starting with this objects allow that the interstinal amountains in objects of societies was diminished after adrenalectomy, had related this defect to a dis toossuus was unmussico arter agrenanceumy man related this octor to a unsure the phosphorylating niechanisms and had assembled rather impressive currance of the phosphory ating mechanisms and non-assemble values impressive evidence that the adrenal cortex was primarily concerned with phosphate transfer Recent attempts to confirm his findings and conclusions have been almost uniform ly unsuccessful (46 47)

DEFICIENCIES RELIEVED BY THE C, STEROIDS

What, then are the primary functions of the adrenal cortex in respect to the what, then are the primary sunctions of the anticial costed in respect to the metabolism of the foodstuffs? The answer appears in those metabolic disturbances in the adrenal ectomized animal which persist despite the maintenance of a normal sodium and potassium balance These include

outin and personner observed these measure

1. Hypoglycemic effect of fairing —Salt treated animals which appear perfectly 1 Myoggyremic egect of Jasting — out tituted annuals which appear perfectly normal and healthy when maintained on an ample diet rapidly deteriorate when food is withdrawn dying in hypoglytema (21, 24, 34) The administration of sugar (in ph) stological saline) rapidly restores them

- 2 Reduced levels of tissue glycogen, particularly that of liver glycogen, during fasting —Thus is due to an mability to manufacture glycogen from the body stores of non-carbohydrate precursors and accounts also for the bypoglycenic effect of fasting (ig. 21, 34, 48)
- 3 Diminished urinary miragen exerction during fasting—In view of the fact that the protein fed adrenalectomized animal excretes normal amounts of nitrogen (21, 49), it seems likely that the difficulty in the fasted adrenalectomized animal is that of mohilization of protein from the tissues and its breakdown to the amino acid stage.
- 4 Disturbance in fat mobilization —Anterior pituitary extracts (50), phlothizm administration (51), or phosphorus poisoning (51) result in the accumulation of fat in the livers of normal animals but fail to do so in the absence of the adrenals
- 5 Alleration of experimental diabetes —The diminution of byperglycemia, gly cosuma and ketosis in depancreatized and phlorhizmized animals which lack the adrenal cortex is readily explained by the disturbances in the mobilization of protein and fat and the consequent dearth of raw materials for gluconeogenesis (19 21.48 52.53)
- 6 Insulin sensitivity —This is not due to the lack of available liver glycogen to combat hypoglycema, for the salt treated adrenalectomized animal with a fauly normal hepatic glycogen level still exhibits the sensitivity (48, 54, 55)
- 7 Muscular weakness -This is alleviated by the administration of carbohy drate (10, 56)

Treatment of fasting adrenalectomized animals with corticosterone or cortin (19 21, 34) restores the normal blood sugar level and, in large doses, may cause hyperglycemia (see Table 31) Such treatment also increases the liver glycogen in normal, as well as in adrenalectomized, animals (19, 21) The muscle glycogen is not so readily affected either by adrenalectomy or by the administration of cortical extracts. Recent work has also confirmed the previous reports that the lack of adrenal cortical hormone diminishes the byperglycemia and glycosuma of diabetes (21, 48, 52) and that the administration of active cortical hormones restores the severity of the diabetic syndrome (28). Sprague et al. (57) have reported a case of a typical diabetes mellitus in a woman which disappeared completely upon the removal of an adtenal cortical timor.

Wells (28) has reported that the injection of phlorhizm into salt treated adread ectomized rats causes them to excrete much smaller amounts of glucose than smaller junceted normal rats. Corticosterone and 17 hydroxy 11-dehydroxorticosterone (Compound E) increase the glucose excretion of the phlorhizmized adrenal ectomized animals to that of phlorhizm treated normal rats. The amorphous fraction (cortin) and desoxycorticosterone have relatively lesser effects (see Table 32).

It may therefore be concluded that the primary metabolic functions of the

admal cortex are concerned with hepatic gluconeogenesis from non-carbohy drate precursors. The observation of Corey and Britton (58) that cortical extracts read the fall of glycogen in perfused livers also suggests an antiglycogenopyte activity of the adrenal cortex. This may explain the more marked effects of cortical extracts on liver glycogen, as compared to muscle glycogen. It also helps to distinguish the action of these extracts from those of the anterior hypophysis (59) (see chap xix, p. 225).

TABLE 3t

LIFECTS OF ADRENALECTOMY AND OF CONTICAL STEROIDS ON THE CARBOHYDRATE LEVELS OF RATS AND MICE (LONG & d) [11])

Cond too	Hormonat The spy	Blood Sugar (Mg per Cent)	Liver Glycogen (per Cent)	Muscle Glycogen (Mg per Cent)
Normal—fed Normal—48 hr fast Normal—48 hr fast	Cortical ex tract	124 80	1 78 0 23 1 64	500 507 536
Adrenalectomy—fed Adrenalectomy—48 hr fast Adrenalectomy—48 hr fast	Cortical ex tract	97 30	2 31 0 07 1 78	533 358 411
Normal—fed Normal—fed	Cortical ex		2 84 9 20	435 2,024
Normal—24 hr fast Normal—24 hr fast	Cortical ex		0 35 8 99	928 223
Normal—24 hr fast Normal—24 hz fast	Corticosterone Dehydrocorti costerone		2 8g 2 25	
Adrenalectomy—14 hr fast Adrenalectomy—14 hr fast Adrenalectomy—24 hr fast	Cortical ex tract		2 28 0 04 2 37	479 258 182
	Normal—fed Normal—45 hr fast Normal—45 hr fast Normal—45 hr fast Adrenalectomy—45 hr fast Adrenalectomy—45 hr fast Normal—fed Normal—16 hr fast Normal—24 hr fast Normal—34 hr fast Normal—34 hr fast Normal—34 hr fast Adrenalectomy—fed Adrenalectomy—fed Adrenalectomy—fe fast	Normal—def Normal—sh frast Nor	Cond to Demonst Cond to Control t	Normal—set Horssonat Segar Casel

MODE OF ACTION OF THE C11 STEROIDS ON CARBOHYDRATE METABOLISM

From their observations on the effect of curtical extract on the R Q of glucosefed adrenalectomized animals, Long Russell, and others (48, 66, 61) have supposed that the adrenal cortical hormone may depress carbohydrate 'ovidation' This conclusion is subject to the usual objections which apply to such use of the R.Q (62) Moreover, Sely e and Dosne (63) have shown that, while cortical extract will inhibit the fall in blood sugar of partially hepatectomized rats, it fails to have any effect in completely liverless animals (confirmed by Remecke [64]) Concordant evidence in patients suffering firm Addison a shease was reported by McBryde and De la Balze (73), who found a very significant increase in the artenovenous blood sugar difference after treatment with cortical extract rich in the C., steroids

- 2 Reduced levels of lissue glycogen, particularly that of liver glycogen, during fasting—This is due to an inability to manufacture glycogen from the hody stores of non carhohydrate precursors and accounts also for the hypoglycemic effect of fasting (10, 21, 34, 48).
- 3 Diminished urinary infrogen excretion during fasting—In view of the fact that the protein fed adrenalectomized animal excretes normal amounts of nitrogen (21, 49), it seems likely that the difficulty in the fasted adrenalectomized animal is that of mobilization of protein from the tissues and its breakdown to the animo and stage.
- 4 Disturbance in fat mobilization —Antenor pituitary extracts (50), phlorhizm administration (51), or phosphorus poisoning (51) result in the accumulation of fat in the livers of normal animals but fail to do so in the absence of the adrenals
- 5 Alleration of experimental diabetes—The diminution of hyperglycemia, gly cosuria and ketosis in depancreatized and phlorhizmized animals which lack the adrenal cortex is readily explained by the disturbances in the mobilization of protein and fat and the consequent dearth of raw materials for gluconeogenesis (19 21, 48, 52, 53)
- 6 Insulin sensitivity —This is not due to the lack of available liver glycogen to combat hypoglycemia, for the salt treated adrenalectomized animal with a fairly normal hepatic glycogen level still exhibits the sensitivity (48, 54, 55)
- 7 Muscular weakness This is alleviated by the administration of carbohy drate (10, 56)

Treatment of fasting adrenalectomized animals with corticosterone or cortin (19 21, 34) restores the normal blood sugar level and, in large doses, may cause hyperglycemia (see Table 31) Such treatment also increases the liver glycogen in normal, as well as in adrenalectomized, animals (19 21) The muscle glycogen is not so readily affected either by adrenalectomy or by the administration of cortical extracts. Recent work has also confirmed the previous reports that the lack of adrenal cortical hormone dimmishes the hyperglycemia and glycosum of dia hetes (21, 48, 52) and that the administration of active cortical hormones restores the severity of the diabetic syndrome (28) Sprague et al. (57) have reported a case of a typical diabetes mellitus in a woman which disappeared completely upon the removal of an adrenal cortical tumor.

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It may therefore be concluded that the primary metabolic functions of the

adrenal cortex are concerned with hepatic gluconeogenesis from non carbohydrate precursors. The observation of Corey and Britton (58) that cortical extracts retard the fall of glycogen in perfused livers also suggests an antiglycogenolytic activity of the adrenal cortex. This may explain the more marked effects of cortical extracts on liver glycogen, as compared to muscle glycogen. It also helps to distinguish the action of these extracts from those of the anterior hypophysis (59) (see chap Xix, p. 225)

TABLE 31

EFFECTS OF ADDENALECTOMY AND OF CONTICAL STEROIDS ON THE CARBOHYDRATE LEVELS OF RATS AND MICE (LONG & d) (217)

Species	Cond tion	Hoymonal Therapy	Blood Sugar (Mg per Ceat)	Liver Glycogen (per Cant)	Muscle Glycogen (Mg per Cent)
Rats	Normal—fed Normal—48 hr fast Normal—48 hr fast	O Cortical ex tract	124 80	1 78 0 23 1 64	590 507 536
	Adrenelectomy—fed Adrenelectomy—48 hr fast Adrenelectomy—48 hr fast	Cortical ex tract	97 30	2 31 0 07 1 78	533 358 411
Mice	Normal—fed Normal—fed	Cortical ex tract		2 84 9 20	435 1,014
	Normal—24 hr fast Normal—24 hr fast	Cortical ex		2 35 2 99	228 228
	Normal—24 hr fast Normal—24 hr fast	Corticosterone Dehydrocorti costerone		2 89 2 26	
	Adrenalectomy—fed Adrenalectomy—24 hr fast Adrenalectomy—24 hr fast	Cortical ex tract		2 18 0 04 2 37	479 158 182

MODE OF ACTION OF THE C11 STEROIDS ON CARBOHYDRATE METABOLISM

From their observations on the effect of cortical extract on the R Q of glucose fed adrenalectomized animals, Long, Russell, and others (48, 60, 61) have supposed that the adrenal cortical hormone may depress carbohydrate "oxidation". This conclusion is subject to the usual objections which apply to such use of the R Q (62) Moreover, Selye and Dosne (63) have shown that, while cortical extract will which the fall in blood sugar of partially bepartectionized rats, it fails to have any effect in completely liverless animals (confirmed by Remecke [64]). Concordant evidence in patients suffering from Addison's disease was reported by McBryde and De la Balze (73), who found a very significant increase in the arteriovenous blood sugar difference after treatment with cortical extract rich in the C., steroids,

despite the fact that this treatment undoubtedly increases the rate of circulation It is apparent, therefore, that cortical extract does not inhibit the uptake of sigar by the peripheral tissue but probably stimulates gluconeogenesis in the liver. It is suggested that its tendency to counteract insulin hypoglycemia (54 55) is exerted in a similar manner.

TABLE 32

EFFECT OF PHIORHIZIN UPON THE EXCRETION OF DEXTROSE AND NITROGENERY RATS UNDER VARVING CONDITIONS OF ENDOCRINE ABLATION AND SUBSTITUTION TREASMY

Endgerine State	Substitution Tree or	DEXTROSE (MG PER 100 GH PER DAY)	NATROORN (BIG PER 100 Gir PER DAY)	pn	COMPARAT VE EXCEE T ON (FER CENT OF NORMAL)	
		(Dext ose	y ttotes
Normal		{62g 574	182 162	3 4	100	100
Adrenal demedulta		624	172	3.6	104	100
Adrenalectomy	NaCt Desoxycorticosterone Corticosterone Compound E Amorphous fract on	142 440 590 (619 560 237	46 124 165 190 155 63	3 7 3 3 3 6 3 3 3 6 3 8	24 74 98 98 40	27 72 93 100 37
Thyroidectomy	l	477	139	3.4	80	81
Adrenalectomy and thyroidectomy	Compound E Compound E+thyroxin	140 382 721	61 103 190	2 3 3 7 3 8	23 64 121	35 60 114
Hypophysectoray	Desoxycotticosterone Corticosterone Compound E Compound E+thyro trophic hormone	148 323 449 412 625	57 100 158 170 196	2 6 3 2 2 8 2 4	25 54 75 69	35 58 93 99

[&]quot;These data a e derived f om the papers of Wells Kendall and amor ates (28 ap 49 73 74)

The probability that the low carbohydrate levels in the fasting adrenalecto mized animal are not due to an increased carbohydrate "oxidation" is enhanced by the demonstration of an impaired work performance of the muscles Ingle(19) has shown that the work performance is markedly dimmished in adrenalectorized animals, even when they are maintained in apparently good condition by a diet high in sodium and low in potassium. This effect is due wholly to the loss of the adrenal cortex for removal of the adrenal medulla has no influence (65). The po

tency of vanous cortical steroids in restoring the ability of the muscles to do work is parallel with their potency as regards carbohydrate metabolism (see Table 3o) Ingle has also shown that the work performance is restored to normal by the administration of glucose in the absence of cortical compounds. These observations would present a curious anomaly if one were to accept the conclusions of Long and co workers as regards the increased 'oxidation' of carbohydrate in adrenalecto nized animals and its suppression by cortical horizones. One would have to reconcide the facts that both the administration of cortical steroids which supposedly suppress glucose "oxidation" and the administration of glucose itself lead to a restoration of normal work performance

The manner in which the adrenal cortex stimulates bepatic gluconcogeness is by no means clear, but evidence is forthcoming that it influences the mobilization and catabolism of both protein and fat Nitrogen exerction is decreased following adrenalectomy, and the administration of cortical extracts restores the introgen output to normal. The increased glycosium observed after the treatment of adre nalectomized depancreatized animals with cortical fractions or steroids is accompanied by a corresponding increase in the urmary introgen. Wells #al. (28) but demonstrated similar effects with the cortical substances in phlorhizmized adrenal extomized rats (Table 32). Another observation which is consistent with the catabolic effect of the adrenal cortects on protein metabolism is that of Fraeskel Conrat #al. (66), who showed that adrenal cortical extracts or the adrenotrophic fraction of the anterior pituitary cause an increase in the level of liver arginase, an enzyme which is concerned in the formation of urea from ammo acida (57).

Concerning the mobilization of fat it had been shown that the phospholipids and latity acids of the blood were decreased following adrenalectiony (68) and that various procedures which increased the fat content of the liver in normal animals usually failed to do so in the absence of the adrenals (69). Barness t al. (43) have recently fed spectroscopically active fatty acids to fasting normal and adrenal ectomized rats. While they were able to identify the administered fat in the livers of their normal animals, this was not the case in the operated animals. The work of Nelson t al. (79) gives an indirect indication of the decreased catabolism of fatty acids after adrenalectomy. They found that the rate of ublization of intra venously injected sodium β by droxybutyrate was markedly reduced in adrenal ectomized rats, as compared to normal animals. Since adrenalectomy does not change the blood ketone level it may be inferred that the production of ketones from fatty acids as diminished in the absence of the adrenals

It should be noted that, while the effect of the adrenal cortex on hepatic glu concepness is unquestionable, there is as yet, little evidence that this influence is a specific one, exerted directly on the liver. The fact that salt treated adrenal ectomized animals, when fed, can maintain good carbohydrate levels suggests that Ur ne N trogen (Mg per Day)

the reduced carbohydrate levels of fasting may result from a disability in the mobilization of protein and fat from the peripheral stores

Finally it should be emphasized that while the separation of adjenal corneal functions into mineral and carbohydrate groups is a convenient point of view there is a certain amount of overlapping of functions. Thus Anderson and Joseph (71) have shown that salt treatment has a beneficial effect upon the fasted adrenal ectomized rat both as regards increasing the survival period during the fast and in

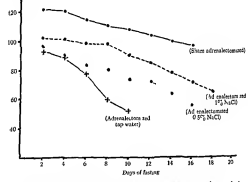


Fig. 51 -- Influence of salt treatment upon the n trogen excretion of the fasted adrenalectomized rat (From the data of Anderson and Joseph 144 711)

creasing the urinary introgen excretion. Figure 51 illustrates their results and in dicates that the maintenance of the mineral halance in adrenalectomized animals does support glucohoegnesis to some extent. A similar slight influence of salt treatment on work performance has also been demonstrated by Ingle (72). It may well be that when all the facts are known the two sets of functions will be found to depend upon the same basic enzyme systems in the cell and that they will be seen to differ only in that each is necessary for a different stage of the reaction chain

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72 INGLE, D J The work performance of adrenalectomized rats maintained on a high sodium 3 . A. P 30L 1 . 73

74

CHAPTER XVIII

THE THYROD

LINICIANS have long recognized the influence of hyper or hypothyroid states on earhohydrate tolerance (t, 2) and on coensting diabetes mel litus in humans (3, 28) in sheep Bodansky (4) found that thyroidectony caused a decrease in the blood sugar level, while thyroxin administration raised it in normal, as well as in thyroidectomized, namnals. However, since thyroidectomy of the normal or depancreatized dog and est apparently has little influence on their earhohydrate tolerance, many writers have been led to minimize the role of the thyroid in this regard (5, 6, 7). It must be pointed out that most of these authors neglected to verify the hypothyroid status of their experimental animals. And since Marine (8) has demonstrated aberrant thyroid issue in over 90 per cent

hormone was administered

METABOLIC EFFECTS OF THYROID HORMOVE

The blood sugar level in hypo or hyperthyroid states is influenced by the effects of the tack or excess of hormone upon the gastro intestinal tract and the

secondary to the changes in metabolic rate, for even large increases in the laturacians day dimitrophenol administration, have no influence on the absorption of carbohydrate. The influence of the thyroid on the rate of absorption of sugar is reflected in the rise and fall of the blood sugar level which follows the ingestion of a carbohydrate meal or the oral administration of sugar solution for festing purposes. In hyporthyroidsm the oral dectrose telerance curve (cf. chap. xx., p. 148) tends to be "diabette" in nature, in hypothyroidsm it tends to be "fist" "The abnormalities are not seen when the factor of intestinal absorption is eliminated by administering the dectrose intravenously.

In the post absorptive state, when the blood sugar is being supplied by the liver, the susceptibility of the latter to glycogenolytic agents or influences has a bearing

* This effect of thyroid is not binuted to the intestinal mucosa but applies also to other epithelial structures eg. kidney tubules (20)

on the blood sugar level. As judged by the results of epinephrin administration, the glycogen in the liver of the hyperthyroid organism is more readily broken down than that in the normal liver. The actual outcome of this state of affairs is, of course, dependent upon the amount of hepatic glycogen present, and this may lead to apparently anomalous results. Thus, Abbott and Van Buskirk (11) have shown that, while the induction of mild hyperthyroidsm leads to an exaggerated hyper glycemic response to epinephrin, severe hyperthyroidism, which depletes the he patic glycogen stores, may lead either to no hyperglycemic response or even to hypogycemia.

2 The glycogen content of itssues other than the liver is also affected by abnormal thyroid states. While lesser degrees of hyperthyroidism have little effect on muscle glycogen, Dambrosi has shown that the administration of large amounts of thy

TABLE 33

RELATION OF VITAMIN B COMPLEX POTHS TO THE EFFECT OF THYROID EXTRACT
ON BODY WEIGHT LAVER WRIGHT, AND LIVER GLYCOGEN
CONTENT (DRILL of of [13])

	BOOM WEIGHT		Lives		TOTAL		
EXPERIMENTAL CONDITIONS	latal (Gm)	Foal (Gm)		Glycogen (per Cent)	Giveogen (Mo.)	Remare	
Control group diet+200 mg yeast Diet 200 mg yeast+100 mg thyroid Diet 200 mg yeast 100 mg thy roid and 1 gm yeast concen trate	215	239	3 5	2 51	86 g	Rats of the same	
	200	161	39	0 34	73 2	used for this work Expen mental period	
	199	208	5 3	2 20	116 T	47 days	

roud hormone definitely interferes with the rate of recovery of glycogen in exer closd muscle (12). Hyperthyrousism also depletes the glycogen of cardiac muscle There is some parallelism between the decreased carbohydrate stores and the in creased exerction of creatine in the time. These effects of the thyroid hormone are not simple in their mechanism, for a lack of the hormone does not produce the opposite results. Hypothyroidism is characterized only by a moderate decrease in the glycogen content of all tissues.

If has become evident recently that the amount of available vitamin B complex has a bearing upon the manifestations of hyperthyroidsm (30)—so much so, indeed, that it will require further work in which the experimental animals or subjects are given ample supplies of vitamin B complex, to demonstrate the pure syndrome of hyperthyroidsm uncomplicated by lack of the vitamin A glumpse of the true picture has been provided in the work of Drill and his co-workers (13), sum arrived in Table 3.3 It may be seen that an amount of vesast concentrate amorem.

mately six times the maintenance dose for normal animals completely counter acted the giy cogen depleting effect of a dose of thyroid which caused very significant loss of glycogen in unprotected animals. It is also important to note that the extra yeast prevented loss in body weight and led to an actual increase in liver weight (3, 14).

- 3 The increased protein catabolism and nitrogen excretion accompanying hyper thyroidism or following the administration of thyroid substances has long been recognized The aggravation of clinical diabetes mellitus by hyperthyroidism and its amelioration in hypothyroid states have linked the thyroid activity on protein breakdown with gluconeogenesis from protein Sternheimer (15) has now shown that the so called "latent period between the injection of thyroxin and the first rise in oxygen consumption is not a period of mactivity. Within 6 hours after the injection of a single dose of thyroxin into rats, he found a loss of liver glycogen and the beginning of a rise in liver protein. These changes became more marked up to about the forty eighth hour and then showed a reversal in direction. By the eighty fourth hour the liver glycogen reached a peak well above the original control level while the total nitrogen of the liver, though falling, was still above the original figures. These and other observations indicated that thyroxin first causes a mobili zation of protein from the peripheral tissues, and also a proliferation of the liver cells, which may be partly at the expense of the initial glycogen stores Subsequently, there is a new formation of carbobydrate from protein Gluconeogenesis from protein has also been observed by Wells et al (16, 17, 18) in phlorhizmized normal, adrenalectomized, and hypophysectomized rats which were treated with thyrona or thyrotrophic hormone (Table 36, D 220)
- 4. In view of the evidence that thyroid hormone stimulates gluconcogenent, it is difficult to understand the relatively minor or negative results as regards carbo-hydrate tolerance which have been obtained either by thyroidectomy of depan creatized animals or by the administration of thyroid substance to such animals In 1938 Dohan and Lukens (19) reinvestigated the effect of thyroidectomy upon

found modification of diabetes which follows removal of the hypophysis from the departereatized animal. However, Sorkin et al. (20) later demonstrated that the administration of thyroxin to hypophysectomized dogs maintained a normal blood sugar level through long periods of fasting and increased their unitary introgen exception to that of fasting normal dogs (Figs. 52 and 53). It is obvious, therefore, that the secondary atrophy of the thyroid gland probably plays an important part in the decreased endogenous protein catabolism and in the related earbohydrate disturbance of the hypophysectomized animal (see chap. xx, 10. 220).

The deficiency in the hypophysectomized animal which is counteracted by the

, це thyroid hormone does not involve the breakdown and transformation of amino acids to sugar, for ingested protein which enters the blood stream as amino acids is readily converted (chap xx, p. 29). The difficulty encountered by the hypoph ysectomized animal during fasting must, therefore, he in the mobilization and breakdown of the body protein to amino acids. It is on this portion of introgen extabolism that the thyroid hormone exerts its influence. This localization of the thyroid hormone effect is supported by certain data obtained in phlorhizin experiments. Lusk and his co-workers (2x xx) showed that fasting thyroidectomized aminals exerted much less sugar and introgen under the mitigence of phlorhizin than

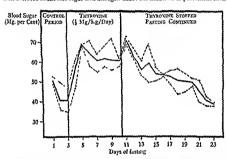


Fig. 23.—Maintenance by thyroune of a normal blood sugar level in a fainting hypophysectomized ong. The upper and lower broken lines, respectively ind cate the maintain and mominum blood sugar levels for each day. The heavy combinuous interinduction the maint washe for all the blood sugar est matering felt least three or days in made on each day. (Swikin and 1 [10])

did similarly treated normal animals. There was no difference between the two types of animals when they were fed protein. Here, again the deficiency arising from the absence of the thyroid was apparently in the mobilization and breakdown of body protein to animo acids.

The question then arises as to why Dohan and Lukena, as well as previous in settigators, were not able to demonstrate the role of the thyroid in depancreatized animals. Indeed, they have recently reported on the subject again [23], this time to the effect that partially depancreatized eats given thyroid extract in doese sufficent to produce tarbycardie and loss of weight did not eithirt any increase in givcosuria Anterior pituitary extract readily increased the sugar excretion in the same animals. We had obtained similar (unpublished) results in our laboratory, not only in depancreatized dogs, but also in depancreatized hypophysectomized (Houssay) animals. One might speculate that the thyroid influences gluconeogene sis from protein in the liver by inhibiting the previously mentioned anabolic action of insulin on protein metabolism. If this were so, thyroid hornone might be expected to have little effect in the absence of the pancreas. But such an action of the thyroid would be difficult to reconcile with the report of Johnston and Maroney.

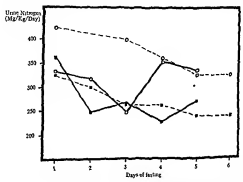


Fig. 33 —Influence of thyrotine on the total ur many nitrogen excretion of hypophysectomized dogs The broken I nee are taken from the figure published by Braier [28] Comparing fasting hypophysectomized

normal dogs

(24) that small amounts of thyroid are anabolic in effect as judged by the positive nutrogen balances obtained in growing children. It would also be out of accord with the evidence that the growth hormone of the anterior pituitary gland is more effective in the presence of the thyroid gland than in its absence and that still greater growth can be obtained when thyroxin is administered along with the growth hormone (25). At the present time, a more hely possibility as regards the

difficulty of demonstrating the gluconeogenetic effect of the thyroid hormone in the absence of the pancreas is that the depancreatized animal given thyroid hormone may become deficient in the vitamin B complex. This is was indicated in the previous section, might prevent the thyroid hormone from producing its characteristic effects.

It should be noted that intensive and long continued treatment with thyroid extract can influence the severity of the diabetic syndrome by damaging the islets of Langerhan (see chap XX, D 243, "McLathyroid Diabetes)

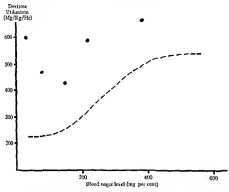


FIG 54—The broken curve represents the utilization of dextrose by normal dogs (see chap xiv p 151) The sold dots represent the sugar utilization by dogs rendered hyperthyro d by the administration of thyronic

5 There is an abnormally rapid rate of carbohydrate itilization by the peripheral itssues of hyperthyroid animals, coincident with the increased amounts of glucose entering the blood from the gastro intestinal tract and from the liver. When thy rown treated dogs are hepatectomized, the rate of fall of the blood sugar level is much greater than in hepatectomized untreated animals (26) Figure 54 compares the actual utilization of carbohydrate of normal and thyroid treated dogs as de

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CHAPTER XIX

THE ANTERIOR PITUITARY

HE relationship between the pituitary gland and carbohydrate metabo lism—diabetes in particular—has been known clinically for a very long time. As early as 1908 Borchardt (1) recognized the large incidence of diabetes in acromegalic patients. American clinicians Goetisch, Usbing, and Jacobson (2, 3, 4) wrote on this subject in 1910, and the relationship continues to be the subject of clinical writing to the present time. It seems certain that, whereas the incidence of diabetes in the general population is about one half of 1 per cent, it occurs among acromegalic patients in about 25-40 per cent of cases Conversely, in hypopituitansm or Simmond's disease, hypoglycemia is often a feature, while Cushing's syndrome, with basophilic adenoma of the pituitary, is often character used by hyperglycemia.

The significance of these clinical observations has now been indicated by the work of physiologists. Curiously enough, the earliest work in this direction was rather misleading, as, for example, when it was found that an extract of the poster nor lobe of the pituitary gland caused a rise in blood sugar level as well as in the blood pressure. More recently, however, the blood sugar raising properties of extracts of the posterior pituitary gland (Pituitrin) have been regarded as being of greater pharmacological than of physiological importance. The remarkable work of the South American physiologist Houssay and of subsequent workers all over the world has shown that it is the anterior lobe of the hypophysis which is important in regard to carbobydrate metabolism.

This relationship was shown by the two chief methods which are the basic proce dures of endocrinologic investigation, namely, the removal of the gland, on the one hand, and the administration of extracts of the gland, on the other The effects of the removal of the antenor lobe of the pituitary gland were first shown by Hous say on toads. The work was later repeated and amplified on dogs, and finally most of the effects have been adequately illustrated by Nature's own experiments on human beings.

The effects of removal of the anterior lobe of the hypophysis in experimental animals or of the destruction of the gland by disease in human beings are as follows

1 Trophic effects — The removal of the pituitary is followed by an atrophy and decreased function of the thyroid gland (5, 6), of the adrenal cortex (7, 8), and of the gonads (0, 10), whether male or female For this reason the pituitary has often

been referred to as "the master gland" of the body. However, the removal of the thyroid or the adrenal cortex or the gonads is followed by histological changes in the pituitary (it?) These changes have been variously interpreted and it is still not quite certain what they mean from a functional standpoint. But there can be no doubt that the removal of these other glands does affect the structure and function of the pituitary. This is also true of the administration of the homomes or extracts of the other glands. Thus, it is clear that, while the pituitary may be more generally important than some of the other glands, it is not merely because it dominates them. It appears rather to co-ordinate the functions of the other glands, so that one might call it "the executive secretary" of the endocrine system rather than the master gland

- 2 A lowering of the blood sugar level —The blood sugar of the hypophysectomized animal under conditions of adequate nutrition is about 20-30 mg per cent lower than the blood sugar of the normal dog (1-, 13 14)
- 3 The hypoglycemic effect of fasting —A normal animal or human being may be fasted indefinitely with little or no effect on the blood sugar level. As a matter of fact, there may be no significant effect until a relatively short time before death from staryation, when the blood sugar may fall precipiously. However in the absence of the hypophysis, fasting is accompanied by rapid development of hypoglycemia so that the animal may die within a relatively short time in hypogly cemic convilsions (12 x x x fo. xr).
- 4 A decreased unine nitrogen exercison (18, 19, 20) —This is due in part to a decreased breakdown of body protein resulting from the secondary thyroid atrophy (see chap xviii, p 214). The atrophy of the adrenal cortex may also be partly responsible (see chap xviii, p 204).
- 5 A decrease in the total metabolism of the body —This is probably accounted for by the depression of thyroid activity, although other factors may be involved. The other factors may be the adrenal cortical atrophy and the loss of weight brought about by the marked anorexia, which is a prominent clinical feature of pututary insufficiency (18, 21, 22).
- 6 An increased sensitivity to insulin —A small amount of insulin which would produce no noticeable effect on a normal animal will, after the removal of the hypophysis cause proloneed and even fatal hypophyciania (r. 2.3. 2.4)
 - 7 A decrease in the potassium content of the blood serum (18, 25)
- 8 A decrease in the reduced glutathione content of blood, liver, and skeletol muscle—The diminished level of reduced glutathione in the liver may be related to the muslin sensitivity (18, 26)
- 9 A cessation of maturation and growth —When the pituitary is removed from immature animals, there is a cessation of maturation and growth (27, 28, 20)

The injection of crude extracts of the anterior lobe of the pituitary into hypoph yectomized animals has been shown to prevent or reverse the consequences of

the removal of the gland Normal animals receiving pituitary extracts exhibit a hypertrophy and hyperfunction of the other endocrine glands (8, 10, 30). Depending upon the conditions, there may be concomitant gain in weight or increased rate of growth, or hyperglycemia, glycosuria, and ketosis may develop (31, 33, 33). Under circumstances in which there is hyperglycemia and glycosuria, there is as an increased excretion of introgen (31, 33). Where gain in weight or an increased rate of growth is a major consequence, there may be a retention of introgen (31, 32).

EXTRACTS OF THE ANTERIOR PITUITARY

The multiplicity of effects resulting from the removal of the gland or the ad ministration of extracts led to many attempts to refine anterior pituitary preparations in such a way as to obtain products with a single or specific activity. Depending upon the method of extraction or purification and upon the test animal and experimental conditions employed, a large number of different anterior pituitary factors have been claimed Collip (34) has recently listed these as follows "growth stimulating thyrotropic, gonadotropic, cotteotropic, lactogenic, diabetogenic, ketogenic, liver fat increasing RQ lowering, blood hind increasing ortgen confunction increasing, anti-insulin, anti-epimephrin, glycotropic, glycostatic, and chromatophore expanding actions"

There are few who believe that these numerous effects obtained under different conditions of experimentation indicate that there are as many separate hormones secreted by the anterior hypophysis Collip suggests that as few as two or three separate hormone proteins may account for all the functional activity. The dosage may play a role, since for example the growth hormone in small doses has only growth effects, while in larger doses it also exerts some corticotrophic and lactogenic action Species differences in the test animals may also be a factor Antenor pituitary extract causes a permanent diabetes in dogs but fails to do so in rats (33) There is also the probability that a number of functions listed by Collip are actual ly duplications of other effects Thus Jensen and Grattan (23) have reported that the anti insulin effect of anterior pituitary extracts is due to the adrenotrophic fraction They found that the administration of adrenotrophic extract, adrenal cortical extract, and corticosterone to mice resulted in a significant resistance to the action of insulin, while the injection of thyrotrophic extract, prolactin, folicie stimulating hormone and thyroxin were without effect Similarly, it has been found that the diminished absorption of glucose by the intestinal tract after hy pophysectomy is probably due to a lack of the thyrotrophic hormone, for it may be corrected by treatment with thyroid hormone (35)

There are also complications of another sort in judging the demonstration of a hormone action when extracts are given or a pland is removed. These complications have to do with the more or less mendental reactions of the entire organism to certain non essential materials contained in the injected gland extracts or to cer

tain secondary reactions of the organism to the condition promoted by the injection of a hormone or the removal of a gland. Thus, Dohan and Lukens (27) have reported that the chronic administration of antenor pituitary extract to depan creatized dogs at first increased and then decreased the severity of the diabetic syndrome. The serum of dogs treated for so months with antenor pituitary extract, when injected into depancreatized animals, reduced their glycosuma and urmary introgen excretion. These results may be likened to the "anti-hormone" effects previously obtained with the gonadotrophic fractions of anterior pituitary extract and, like them, are probably due to non specific antibodies formed in response to the proteins contained in the impected extract.

The decreased food intake which leads to marked undernutrition following hypophysectomy may also be responsible for some of the results usually attributed specifically to the lack of the pituitary hormones Mulmos and Pomerantz (18) studied the effects in rats of complete manition during starvation and of chimnic undernutration resulting from an allowance of approximately half the normal food intake. They found that the loss of weight and the histological changes in the endo crine glands resembled those following hypophysectomy. The authors concluded that manition affected the anterior hypophysis in such a manner as to reduce its secretion of the timphic hormones. It would be interesting to know whether all their results would or would not have been prevented by the injection of anterior pituitary extracts into their chronically undernourished animals. Levin (30) has recently shown that the decrease in weight of the viscera, which follows hypophy sectomy, can be completely prevented by force feeding the animals to the level of normal food intake. Since such treatment, however, does not restore the weights of the endocrine glands their atrophy is linked directly to the loss of the trophic hormones

In view of the attendant difficulties it is not surprising that the results of at tempts at the separation and punification of the vanous fractions of antenor pittu tary extracts continue to be difficult to harmonize and continue to disclose hitherto unsuspected effects. Bergman et al. (40) believe they have separated four entities from antenor pittutary extracts—namely, lactogenic, thyrotophic, gonadotrophic, and the carbohydrate metabolism factor. Meaniber et al. (41) have reported that preceptiation with cysteue enabled them to separate the lactogenic and thy rotrophic effects from growth fractions of antenor patientary extract, this procedure resulting in the preparation of almost pure growth horizone. Graves and his convokers (42) have described the properties of a more purified diabetogenic factor extracted at pH 11 It was non-dialyzable and was destroyed by a temperature of cot of C for 12 minutes at pH 10. This diabetogenic fareful was ketogenic and lowered the R Q It was rich in the growth factor but exhibited hitle prolactin action.

Teague (43) has reinvestigated the association of the melanophore hormone

with the "specific metabolic principle of the pituitary" previously reported by Collip and his co-workers (34, 44, 45) According to Teague, preparations of the pituitary gland rich in melanophore hormone, obtained from various sources and prepared by different methods, varied considerably in their effect on oxygen con sumption in rats. The melanophore activity of extracts could be selectively de stroyed without removing the metabolic effects. It was concluded that the melano phore hormone was not identical with a substance in the pituitary extracts which would increase the metaholic rate. It was further pointed out that the results did not support the existence of a specific metabolic principle of the hypophysis, since it was found, in the course of the work, that metabolic stimulation was produced by a pituitary extract after treatment with acid and after tryptic digestion, and since such metabolic responses were occasionally obtained with extracts of muscle, liver, and kidney Collip (34, 45) has also reported the action of a pituitary ex tract which stimulates the "dark" cells of the adrenal medulla without affecting the chromaffin tissue. The extract is active when administered by mouth. The sig nificance of this action must await enlightenment as to the function of the "dark" cells Finally, Houchin (46) has been able to decrease the alkali soluble protein components of the liver with anterior pituitary extract fractions and has suggested the existence of a protein metabolism hormone which is distinct from the lacto genic, thyrotrophic, carbobydrate metabolism, fat metabolism, and gonadotroph ic hormones

Probably the hest isolation of purified anterior pituitary hormones from the standpoint of methodology and their most accurate characterization from the biological standpoint are to be found in the work of Fraenkel Courat et al (47)

practically all the known metabolic effects of crude extracts of anterior pituitally are accounted for, except the ketogenic There is, at present, no way of rationalizing the distribution of the various effects among the different hormonal entities, nor is it possible to say whether or not some of the effects obtained with the growth and lactogenic hormones are mediated by one or more of the endocrine glands. Furthermore, the separation of practically pure entities still does not preclude the possibility that they are fragments of a single complex original hormone. It is not yous that much work remains to be done in this field.

THE INFLUENCE OF THE ANIERIOR PITUITARY AS A WHOLE ON VARIOUS ASPECTS OF CARBOHYDRATE METABOLISM

The well fed hypophysectomized animal maintains a significantly lower blood

sugar level as will be discussed later (chap xx1 p 255) The profound influence of sugar sever as with the carbohydrate levels of blood and tissues is most clear. the anterior puturisary on the carbonyurate severs of misord and tresuces to into a tree.

ly demonstrated by observing the effects of fasting. When food is withheld from 225 by demonstrated by observing the enects of fasting. When food is withheld from the hypophysectomized organism, there occurs a progressive drop in the blood the hypophysic totalized organism there occurs a progressive drop in the mood sugar level terminating in hypoglycemic convulsions and death (12-13). The gly sugar level terminating in nypogitycenuc convilisions and ucasin (12 (3) 1 inc 80) cogen content of the fusices is decreased particularly that of the liver (12 40 50) cogen content of the tissues is decreased particularly that of the diver (12, 49, 50). This occurs even when the pancreas and the hypophysis are both removed (12, 49, 50). run occurs even when the painters and the hypophysis are both removed (12) and the effect of fasting is exaggerated by the administration of phlorhizin (12 15)

METABOLIC ACTIONS OF PURIFIED ANTERIOR PITUITARY HORMONES

	TICTIONS OF Prince		
Rarmone	AG	UOR Draw	
Grand A		TOP YEAR HOP	Man -
Growth (GH)	Act ros		AU LES
	N trogen retention (a present	1	THE REAL PROPERTY.
	adequate Petention Communication	Remarks	
	a Increase insulin)	cof Gir	Refe ences
		DOTA TOTAL	on //
	ly departreatized an mals	at Pose each oth	as US 47 74
	S Increase of ment in pancrese	and Tire.	H
drenotrophic (ACTH)	S Increase of insulin in pancreas Decrease of insulin in blood becrease in liver arginase	ergustically sy	ra /
(MC11D)		1	1
	2 Increase a a tracen	1	- 1
	2 Increase n i ver argunase	Ivan	1
Thyroteen	3 Inhib tion of insuhn action 4 Increase in liver above	V a adrenal cortex	1.
Thyrotrophic (TII)	Increase in liver glycogen	1	(47 48 74)
•		1	1
	2 Increase n besal metabol c rate	I	1
I.a.		Nos t a mud .	1
Lactogenic (LII)	3 Increase in basal metabol c rate 4 Decrease in trisue NPN	Nos t 2 and 3 viz	(47 77)
	1 1		1 " "
	Z Description of neutron of		1
	I Increase of asulm a pancreas Decrease of insul n a blood		l
These -	12 Diood		
ant effects of fact.		-	(74)
anterior pituitary exert	ng might be interpreted		
Isone- Wall Creve.	O. O. DILLETTON . 1	-	

These effects of fasting might be interpreted in one of two ways either (1) the ances energy or tasting august of interpreted in one of two ways entire (1) the antended partially exerts an inhibitory influence on carbohydrate utilization by the automy printerly exerts an infimitivy infinitely consequence on entonyment attraction by in-lesses and hence hypophysectomy is followed by an excessive rate of utilization mith which the capacity of the liver for gluconeogenesis cannot keep pace or (2) the gland exerts its primary influence on gluconeogenetic processes in the liver see, seems exerts its primary innuence on governesseems processes in the average and hence its removal leads to a reduced rate of sugar formation from non-carboand mence its removal leads to a requeed rate of sugar necessary even for normal utilization can no longer be supplied. It is clear that these alternative explanations remains the catent that they depend upon a disproportion between the rates as a minar to the extent that they depend upon a disproportion occurred the fact of Sugar formation and sugar utilization. But the first explanation attributes the Dut the miss expandation and sugar attribution for the miss expandation attributes the second attributes it to the It the present time the evidence that is available regarding the foresome av

hepatic gluconeogenesis in intact non anesthetized normal and hypophysectomized animals by means of the London cannula technic. From the blood sugar on tents of the inflowing and outflowing hepatic blood, they estimated that the rate of sugar output from the livers of their fasting hypophysectomized dogs was only about 50 per cent of the output from the livers of fasting normal dogs. The work of Wells and others (60 of) in Kendall's laboratory confirmed the defect in gluconeogenesis in hypophysectomized animals and indicated that this influence of the hypophysis was exerted partly through the adrenal cortex and partly through the thyroid gland. These workers studied the urmary sugar and introgen exception of normal adrenalectomized thyroidectomized and hypophysectomized aris in spectively treated with phlorhizin. They also included animals from which both

Table 35°

Relative Stability of Muscle Glycogen after Hypophysectomy
(Sorem Levine and Lemmn [48])

Compte on	No or Docs	(Mg and Cities)		AV BLOOD LACTIC ACTS (Ma 722 CENT		AT DE- CHEASE DI MUSCLE GLYCOGEN	AV IN- CEXAME IN BLOOD LACTIC ACID
		In tal	Fral	In tal	Fnel	(Ma res Cent pre He)	(Mares Cent yea Ha)
Normal Normal given an	15	511	355	50 5	106 7	43 [‡]	15 1
terior p tuitary	32	601	448	1 40 0	1246	42 5	182
Departreatized Hypophysecto	13	337	217	50 0 118 T	183 2	42 S 38 4	31.0
mized	5	584	570	27 3	62 8	4 2	9.5

Observe a mustle glycogen and in blood factor and in I we less dogs during experiments for which the blood sugar was mand at/or above the aurum level by constant intertion of glucore.

the thyroid and the adrenal glands had been removed. By administering various hormones and combinations of hormones to the operated rats they were able to judge which hormonal factors restored the hypophysectomized animals to a nor mal response so far as sugar and integen exerction were concerned. Their results are summarized in Table 36. It may be seen that neither thyroid nor adrenal cortical hormone by itself was able to rectify the deficiency in hypophysectomized rashwhile the combination of both hormones was successful. It may be concluded that the gluconeogenetic influence of the thyroid gland (chap. xviii) and of the adrenal corticx (chap. xviii) are each partly responsible for the total effect of the anterior initiative.

INFLUENCE OF THE ANTERIOR PITUITARY ON CLUCONEOGENESIS FROM PROTEIN AND FAT

Figure 56 compares the effects of exclusive fat or protein feeding and of fasting on the blood sugar level of a hypophysectomized dog. It may be seen that the am

mal has no difficulty in maintaining its blood sugar level at the expense of inmay has no connectively in maintaining its blood sugar level at the expense of in-gested protein. It cannot maintain this level when it receives only fat. It is also gested protein at cannot manufaut the level which it receives only lat at is also evident that the length of time which the animal can withstand fasting depends evident that the length of time which the summar can withstand lasting depends upon its previous feedings. After a protein feeding period of to days it took about 220 upon its previous recuings after a protein recuing period of 10 days it floor about 12 days of fasting to reduce the blood sugar to a consistently severe hypoglycemic 12 days or tasting to requee the blood sugar to a consistently severe hypogycemic evel, after a prolonged fasting period and a rapid recovery of the blood sugar evel, after a protonged tasting period and a rapid recovery of the notice sugar evel by the administration of protein for 1 day, a second fasting period resulted

The most obvious explanation for the ease with which the hypophysectomized the most ouvrous explanation for the case with winter the hypothysectionized intelligence or maintain its blood significant from protein placed in the may can restore or maintain its mood sugar sever from protein praced in the its unable to utilize adequately the much

Effects of Endocrine States and Substitution Therapy ON PHIORHIZIN DIABETES IN THE EATS Anna Anna Anna Anti-

	A STATE OF THE PARTY OF THE PAR	V DIABETES	IN THE RA	YOU TO-		
CONDITION OF ANIMALS		-	THE RA	I.	KAPY	
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larger amount of its own tissue protein for the same purpose, is the fact that in easts amount or its own usue protein for the same purpose, is the fact that in Rested protein enters the blood stream as amino acid. It may be concluded that seases proven enters the blood stream as anniho acad at may we concluded that the antenor pituitary exerts its influence on gluconeogeness from protein by faciliand the conversion of the breakdown of tassue proteins to the amino acid stage aring the conversion of the oreaktown of tissue proteins to the amino acid stage of the influence of previous protein feeding on the hypoglycemia of fasting is suggests that the anterior pituitary may control proteolytic processes within see suggests that the antenior pituntary may control protective processes within the cells but not be important for the transport and conversion of so-called over the put not be important for the transport and conversion or so-caused storage, protein (78). The influence of the anterior puturary on the breakdown swage protein (76) The innuence of the anterior parametry on the oreakoown of protein to amino acids is exerted—in part at feast—through the thyroid gland on protein to amino acids is exerted—in part as reast—tunougo the trythologistic lines has been shown by experiments in which the blood sugar level of fasting hus ness neen snown ny experiments in bouch the onord sugar sever or rusting the physicismized dogs has been maintained indefinitely by the administration "Prophysectomized dogs has been maintained indemnitely by the administration of thyroin (Fig. 52, p. 215). The thyroun simultaneously restores the nitrogen excretion of these animals to that of fasting normal dogs (20) That there is a difficulty in gluconeogenesis from the fat stores of the hypophy

sectomized animal is evident from the fact that fasting may induce a fatal hypoglycema even though ample deposits of adipose tissue are present. The influence of anterior pitutary extracts on gluconeogenesis from endogenous fat in normal animals was shown by the work of Neuteld, Scoggan, and Stewart (62). They m jected various anterior pituitary extracts as prepared in Collip's laboratory, into female mice and made chemical determinations of the entire carcasses of their aminals. They found an increase in the total glycogen content, a decrease in the amount of fatty acids present, and no change in the introgen. The inability of the hypophysectomized animal to maintain its blood sugar at the expense of ingested fat may depend upon the fact that this foodstuff is absorbed into the blood in the

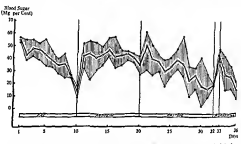


Fig. 56—Effect of exclusive fat or protein feeding and of fasting on the blood sugar level of the hypophysectomized dog. The shaded area represents the spread of the blood sugar values and was obtained as the spread of the blood sugar values and was obtained as the blood of the spread of o

[13])

nc na + al fa+ Tinlike ingested hypophy

THE HYPOPHYSECTOMIZED-DEPANCREATIZED (HOUSSAY) ANIMAL

In 1930 Houssay and his associates reported their observations on hypophysec tomized depancreatized dogs (12, 15). They found that such animals exhibited less severe diabetes than dogs with only the pancreas removed. The blood sugar level varied in different animals from 320 to 113 mg per cent. Sometimes spontaneous

THE ANTERIOR PITUITARY hypoglycemia occurred The glycosuria was correspondingly variable and was en hypogycemia occurred and gyrusuna was currespondingly variable and was on tirely absent in some cases. Natiogen excretion was only slightly decreased but they also the very mild or absent. The animals survived for months without 221 msulm

suun Figure 57 shows that fasting has the same hypoglycemic effect on the Houssay Figure 57 snows that rasting has the same hypoglycenine eners on the hypophysectomized animal (13). It also indicates the quantita ong as it has on the hypophysection and a triple it also includes the quantitative relationship between the amount of protein ingested and the consequent rise and the blood sugar level As might be expected the glycosum also depends upon (Me per Cent)

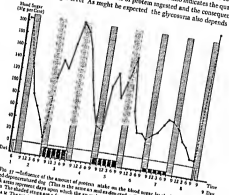


Fig 37—Loftmence of the amount of protein naide on the blood topic level of the hypophysical Fig. 57—Influence of the amount of protein make on the blood sugar level of the hypophysical confidence of the amount of protein make on the blood sugar level of the hypophysical of the superstanding of the superstandin to depondent to do (This is the same as males do cled in F.E. 50 before parameterizing). The class that arrows appear which the so and was fed the superimposed what a tross indicate the cal size represent days upon which the 20 mal was fed the superimposed white arrows indicate the size of the 10 mal. The bladed stops are a foreshortened representation of the n gat periods. Lettering 0.0 FM and 0.0 mal. The bladed stops are a foreshortened representation of the n gat periods. meat The shaded steps are a fore-horizoned representation of the grad periods between 9 00 P.M. and 90 0.0 M. The folds amount of food given on the representation of steps and forest days of feeding was a followed day; 4.00 km of measurements above the steps forest days of forest forest forest days of forest forest forest days of f 9 oo as. The folds amount of food given on the respective days of feeding was as follows: day 5 400 cm of clean mast, form of cance sage; 130 cm of the pancress day 3 378 gm of protecting learn goest, of the pancress day 3 378 gm of protecting learn goest, of the pancress day 3 378 gm of protecting learn goest, of the pancress day 3 378 gm of pancress days 3 400 gm. of teat meat to gen of case sugar 130 gen of saw pancress chy J 378 gen of protein as least meat.
1505 gen of protein chay 7 00 gen of protein a evening of chy 2 same as chy 2 (Soat act of [13])

the protein intake as is shown in Table 37. It should be noted that regardless of one process maske as is snown in 1400# 37 at should be noted that regardless of the degree of diabetic manifestations in the different animals no ketonina nas observed

If may be concluded that the same disturbance which causes the disability of he hypophysectomized animal as regards maintenance of his blood sugar level, namely the impairment in gluconcogenesis is also responsible for the ameliora

tion of diabetes in the Houssay animal. The extreme variability in the seventy of the diabetic syndrome noted by Houssay and other authors undoubtedly resulted from the variability of the food intake of their experimental animals. The well fed Houssay animal actually exhibits a diabetic syndrome of moderate severity, except for the lack of ketosis. The undernourished Houssay animal manifests little or no diabetes. But even under the most favorable nutritional conditions, the diabetic syndrome is not as intense as in the depancreatized animal with the hypophysis intact. This is readily understood when one considers the unavail

TABLE 37

HYPOPHYSECTOMIZED DEPANCREATIZED DOGS (SOSKIN et al. [141])

Dog	Survival (# thout Ipsul n) (Weeks)	D et (400 Gm Mest 60 Gm Sugar 220 Gm Pantreas)	Ketonuria	Average Glucose Excret on (Gin per 34 H s)	Average N trogen Excret on (Gm per 24 Hrs.)	Aversão D N Ratio
H 7	4	Full Partial (4)	None None	10 E 2 4	5 t	÷
H 21	6	Full	None	800	14 3	14
H 35	7	Full Parcial (3)	None None	75 °	11 7 4 9	1 28 0
H 74	9	Full Partial (4)	None None	83 o 33 5	75 9	1 50 0 50
Н за	23	Full Partial (o)	None None	70 3	12 0	o 86
Sally	74	Fuli Parnal (3)	None None	6r 8	15 0 14 0	0 12 0
H4	15	Full Partial (1)	None None	95 9	16 5	2 10 2 50

^{*} Th , was calculated after subtract ug the amount of sugar ingested f on the glace e excreted

ability of its endogenous protein and fat for gluconeogenesis and the fact that, of the ingested food materials, only sugar (as such) or protein (amino acids) can con tribute to the maintenance of the blood sugar level. In other words, while the depanceatized animal with hypophysis intact can make excessive sugar at the expense of both protein and fat (endogenous or exogenous), the Houssay animal can use only ingested protein for this purpose. This accounts for the hypoglycemic effects of fasting, in spate of ample fat stores, the low D. N. ratios, the lack of keto six, and the relatively long survival without insulin.

The amelioration of the diabetic syndrome in the absence of the hypophysis re sembles, in many respects, that seen in deparcreatized dogs maintained without insulin on undernutrition diets composed solely of protein (63–64) It has been

shown that carboby drate utilization proceeds at a normal rate in untreated pan soom taat carooyunate umnaaton proceess at a normal rate in uniteaten pan creatic diabetes (chap xvi, p 185) and that hypophysectomy decreases carbohycrease canceres tenap xvi, p 1053 and that hypophysectomy decreases carbohydrate utilization (Fig. 55). Hence, neither undernutration nor hypophysectomy orate utuazation (1 ig 55) mence, neutner undernutation nor hypophysectomy can be held to amediorate the diabetic syndrome by restoring carbohy drate utubiza 233 can be near to amenorate the madetic syndrome by restoring carbohy drate utilization. Undernourished depancreatized animals survive from 4 to 6 weeks and, de ton Undernourished departeratized animals survive from 4 to 0 weeks and, de spite the complete absence of insulin, become progressively less diabetic the longer spite the complete assence or maum, become progressively less unabetic the longer they survive. There is a progressive lowering of the D. N ratio, a gradual increase. tary survive there is a progressive sowering or the D IV ratio, a gradual increase in the RQ, and an increasing retention of administered sugar which has both proin the κ V, and an increasing retention of administered sugar which has both protein spanning and anti-togenic actions. These criteria of "carbohydrate oudation" ten sparing and antiketogenic actions. Liese criteria of "carbonydrate ordation" become apparent as the fat stores of the animals are depleted. The difference be occome apparent as the tat stores of the animals are depicted the difference be theren these animals and Houssay dogs consists in the means by which the dia theen these animais and moussay nogs consists in the means my which the dia better is modified rather than in any difference in the final state which is reached octes is moduled rather than in any difference in the linal state volico is reached. The undernourished departrealized animals suffer a gradual and incomplete loss and undernournance departmentized anumans some a gradual and incomplete loss of body fat as the period of undernutrition progresses, while the Houssay animals or body rat as the period of undermutrition progresses, while the rioussay animals ethibit an acute loss of ability to utilize the ample fat stores which are present In cunter an acute loss of admity to utilize the ample that stores which are present in both cases this leads to a decreased new sugar formation, so that utilization of carbohydrate is unmasked

INFLUENCE OF THE ANTERIOR PITUITARY ON

The mechanism of the increased sensitivity of hypophysectomized animals to THE INTERIOR OF LINE INCREASED SEISHIVITY OF MYPOPHYSECTORINGED ARRIVAGE INBUILD IS NOT COMPLETEly understood. It may depend on any or all of the following insum is not completely understood at may depend on any of an or the following factors (a) a lack of counterregulatory response to hypoglycema by the liver of tectors (e) a tack of counterregulatory response to hypogrycemia by the inver of the bypophysectomized animal (b) a decreased rate of inactivation of insulin by the blood and tassues of the bypophysectomized animal, so that the administra tion of a given dose of insulin might result in the presence of much larger effective quantities of the hormone, and (c) the absence in the bypophysectomized animal quantities of the normone, and tell the absence in the oppopulationized animal of an anti-insulin factor which antagonizes the action of insulin in the extrahepatic

The decreased rate of gluconcogenesis in the liver of the hypophysectomized Annal may be a factor which limits the ability of the animal to restore its blood auman may be a factor which aims sine ammy or the amman to restore its blood signar level. This agrees with the fact that adrenotrophic hormone or adrenal corti ougast revet 1 ms agrees with the fact that agreenorophus normous or agreem cortical extracts which increase bepatic gluconeogenesis also restore the normal recat extracts which increase nepatic gluconeogenesis also restore the normal re-sponse to insulin (23). But gluconeogenesis cannot be the only factor, because thysponse to mainin (23) that gluconeogenesis cannot be the only factor, because thy-form, which resembles adrenal cortical extract in increasing gluconeogenesis, does not affect the insulin hypersensitivity of hypophysectomized animals (Fig. 58)

or americ time insuum hypersensitivity of hypophysectomized animais (Fig. 58). The work of kepinov (65) and that of Bodo (14, 66) indicate that the suscepti hire work or repinov (03) and mar or more (14), one manuale that the susception of hyer glycogen to breakdown by epinephini is diminished in the absence of only on a ver grycegen to preakuown by epinepinin is ununissed in the absence of the pyophy sis. If this applies to the endogenous secretion of the adrenal mediulnormally evoked by hypoglycema, it would, of course, contralsed after dynamics.

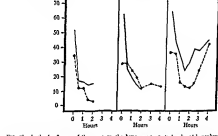
Blood Sugar

(Mg per Cent)

It seems likely that the mactivation of insulin in the body is accomplished by sulphydryl compounds (67, 68). It has been shown that muscle extracts mactivate insulin in vitro by virtue of two components, one of which is probably reduced glutathione (GSH), while the other is the SH groups of proteins (68). The application of these facts to the intact hiving organism is indicated by the observation that the intravenous administration of cysteme is followed by decreased sensitivity to insulin. It has also been shown that the livers of hypophysectomized rats have a significantly lower GSH content than those of normal rats (26). There is, therefore, some basis for supposing that the increased effect of insulin in the absence of

√r unit/kg

d unit/kg



√r unit/kg

Fig. 58—Lack of influence of thyroxine on the hypersenaturity to Insulin of hypophysicionized dogs Each set of curves represents a different hypophysicionized named. In each case the broken his represents the effect of insulin before thyrozine treatment while the continuous his represents the effect of the same dose of insulin during thyrozine administration. Note the higher init al blood sugar values in the thyroxine treated animals. Sockint of # 100.

the hypophysis may be due to a prolonged period of action because of a decreased rate of inactivation

The work of Himsworth (69) may be taken to indicate the presence of a periph eral anti insulin factor in the hypophysis. He reported that while the administration of crude printiary extracts did not influence the spontaneous fall of the blood sugar in hepatectomized rabbits it did interfere with the accelerating effect of in sulin upon the rate of fall. The results of Russell et al. (see p. 226) appear to support Himsworth's observation. But the evidence of both is opposed by the find

ings of others (p 226) which are incompatible with the conclusion that the ante nor pituitary exerts an important pempheral action or primitary exerts an important peripaeral action.

It is evident that the sensitivity to insulin of the hypophysectomized animal de 235

At is evident to at the sensitivity to insum or the appoposysectomized animal de pends—partly at least—on the liver Whether or not there is a peripheral factor pengs—parity at seast—on the over whether or not there is a peripoeral factor in the sensitivity must await further work. The use of the more recently available in the sensitivity must await further work—the use of the more recently available pure trophic fractions of the anterior pituitary in the liverless animal should make INTERDEPENDENCE OF THE METABOLIC FACTORS

Table 34 summanzes the various known physiologic effects of the best isolated same 34 summarises the various known physiologic enerts of for pest isolated components of anterior pituitary extract which together exert the so-called dia touspussents or anterior pituntary extract which together exert the so-caused dia betogenic action. It will be noted that the most important factors are the adrenooctogenic action at will be noted that the most important factors are the adreno-trophic thyrotrophic and growth hormones. In general, these hormones act hy motopine invroteganc and grown normones in general these normones act by mo-bilizing the non-carbohydrate precursors of blood sugar from the periphery and by ontaing the non-carnonyurate precursors or brood sogar from the perspirery and by stimulating gluconeogeness at their expense in the liver. This seems an anomalous summating graconeogenesis at their expense in the liver. Anis seems an anomalous function to attribute to the growth hormone, since the process of growth must in volve protein synthesis and attrogen retention rather than the reverse The fact is voire protein syntoesis and introgen retention rather than the reverse. The fact is that the growth bormone exhibits either its anabol c or its catabolic action de tuat the Browsta optimine exhibits either its analysis of its catabout action the pending upon the presence of absence of insulin (33, 70, 71). In the normal animal penantig upon the presence of absence of insulin 133.70-71). In the normal animal of in the departreatized animal receiving large amounts of insulin the growth bor we make dependence animal receiving large amounts of insum the growth bor mone causes nitrogen retention. In the untreated diabetic animal it causes in Certain experiments showing the amelioration of the diabetic syndrome by

certain experiments snowing the amenioration of the mapping symmotion of advended animal by the earning-tumy and its exaceroation even in the hypophysecunitized animal by the administration of large amounts of adrenal cortical hormone have been interpreted aum martation or targe amounts of auternal corried normone have oven interpreted as indicating that the adrenotrophic hormone is the most important factor in the as muscating that the antenotrophic normole is the lower important factor in the dashedgenic action of the antenor pituitary (72-73). This is not necessarily so It outsetsgeene action of the anterior pituitary (72-73). Links is not necessarily so it is true that the presence of some adrenal cortical hormone is essential for the dia a stee that the presence of some agrenal cortical normone is essential for the dia betogenic action of the other antenor intuitary factors and this may account for occuprance action of the other antenor partitions factors and this may account for the amelioration of diabetes in its absence. But it has also been shown that the the annunation of diabetes in its absence but it has another shown that the administration to an adrenal ectomized an mail of an amount of adrenal cortical autumatration to an aurenaiccionizeu an mai oi an amount oi aurenai cortical horizone which by itself exerts no obvious diabetogenic effect wil enable that am nonmore which my steel exerts no obvious manetogenic chief will enable that and to yield a significant diabetogenic response to anterior pituitary extracts

The situation is probably not so complicated as appears at present. If we reare situation is propany not so computated as appears at present, it we re gard each of the hormones as a catalytic influence at a different point in the chain serve cast, or the normones as a catalytic numerice at a numeric point in the chain of reactions responsible for the mobilization and catabolism of the foodstuffs at executions responsible for the monumentum and cambusium of the foodsitude it student that the acceleration of any one of the reactions may increase the rate o evigent that the acceleration of any one of the reactions may increase the rate of the whole chain. However, the absence of any one of the hormones may lead to on the project chain frowever the absence of any one or the normones may read to such a bottleneck at its particular point of action that the accelerating effect of any or all of the hormones may be nullified

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CHAPTER XX

PERMANENT EXPERIMENTAL DIABETES PRO DUCED WITHOUT SURGERY

HE diabetic syndrome induced in certain laboratory animals during the injection of anterior pituitary extracts may be termed 'hypophyseal (or pituitary) diabetes. As first shown by Evans (t) and by Houssay and his co workers (2) and subsequently confirmed by many others this type of experimental diabetes begins to diminish in intensity after a few days even while the injections of extract are continued. The syndrome disappears very rapidly following the cessation of treatment (3)

In 1017 Young (4) reported that the injection of increasing massive doses of crude antenor nituitary extracts into dogs resulted in some animals in a nerma nent diabetes which persisted indefinitely after the injections were stopped. He also reported experiments in species other than the dog. He found that the mouse rat and guinea pig showed hardly any effect from the injection of his crude ante nor pituitary extract. About half of the rabbits and rats showed slight and transi tory diabetogenic effects. Very young dogs or punpies resembled the rabbit and cat rather than the adult dog (5 6 7 8) Lukens and Dohan (6) were able to dem onstrate the diabetogenic action of nituitary extracts and the production of perma nent diabetes in partially deparcreatized cats Richardson (10) made histological studies of the pancreatic glands of dogs rendered permanently diabetic with pitui tary extracts and reported that the islets exhibit reduction in size, hyalinization and degranulation of the B-cells Best et al (11) found that the pancreas of such does contains from 0 to 0.2 units of insulin per cram, as compared with the average figure of 3 4 units per gram in the normal animal. The fact that dogs can be ren dered permanently diabetic with anterior pituitary extract but that this is not possible in rats may be explained in part by the observations of Marks and loung (12) They confirmed the decrease in the pancreatic insulin content in the dog but found that the administration of anterior pituitary extract to rats in creased the amount of insulin in the pancreas. They reported that in this respect the rabbit behaved like the dog while the mouse resembled the rat

It is important to distinguish between the experimental diabetes seen during the injection of hypophyseal extracts (before the destruction of the islets of Langer ham and reversible) and the perimanent diabetes which persists after cessation of antenor pituitary injections and which is not very different from pancreatic diabetes Hence, it seems wise to adopt the nomenclature suggested by Houssy (13)

and to reserve the term "hypophyscal (or pituitary) diabetes" for the temporary state during hypophyscal injections, while using the term "metahypophyscal diabetes" for the permanent syndrome resulting from the destruction of islet tissue

METAHYPOPHYSEAL (YOUNG'S) DIABETES

Despite their fundamental similarity, there are certain, as yet unexplained, differences between the metabolism of depancreatized dogs and that of dogs with metahypophyseal diabetes. We quote Marks and Young's own summary (14) of their findings and conclusions regarding the latter type of animal. These authors used the term "pituitary diabetic" to denote the metahypophyseal syndrome

- r Dogs made permanently diabetic by treatment with anterior extract differ most obviously from depancreatized dogs in the following respects
 - a) Some of these dogs require more insulin for the control of glycosuma than do depan creatized dogs.
 - b) The pituitary diabetic dogs are able to survive for long periods in good health without in sulin therapy, if sufficient utilizable food is given. The intensity of the diabetic condition may vary from animal to animal.
- 2 Removal of the pancreas from a pituitary diabetic dog resulted in a slight and possibly not significant fall in insulin requirement. The pancreas contained 2 g units of insulin, compared with an average figure for nine normal dogs, of comparable weight, of 76 units.
- 3 On a protein diet, the pituitary-diabetic dogs exhibited hyperglycemia, a substantial glyon suria and ketonura, with a D/N quotient of over 3 o in most instances, on a high drate diet, these dogs returned about 5.9 per cent of the total available carbohydrate in the food, on a diet of beef suct, the blood sugar level the glyosuma and ketonura of these dogs.

sulted in a substantial rise in ketonuria. These results support the conclusions of Petral (1924), which were drawn from clinical investigations that protein (meat food), and not fat, is particularly concerned in the actiology of ketonuria.

- 4 The metabolic rate of the pituitary-diabetic dogs was somewhat above that of control nor mail animal under similar conditions, but the excess above normal was not so great as was found with depanceatized dogs
- 5 As indicated by the hypoglycemic effectiveness of 5 units of injected insulin, by the Hims worth (1036) glucose insulin test, and by the de Wesselow Griffiths (1036) serum test, the pitutary-diabetic dogs do not possess any abnormal degree of insulin insensitivity.

The state of the s

There are two additional items in their paper, not mentioned in the summary, which seem of particular interest. In following up their observation of the ketogenic effect of raw meat, as compared with casein in their "pituitary-diabetic" amals, they found that the residue of raw meat which had been repeatedly extracted.

^{*}This statement applies only to metahypophyseal diabetes. In hypophyseal diabetes there is a marked insensitivity to insulin (8)

with hot water exerted only about one-quarter of the ketogenic effect exerted by the original amount of the raw meat. The supplementation of the extracted meat with a concentrate of the hot aqueous extract caused a significant increase in ketonuria. Marks and Young also made a number of comparisons between their results and those obtained by Langfeldt (13) on partially depancreatized animals one might speculate as to the extent to which the differences between meta hypophyseal diahetes and pancreatic diabetes might be caused by the presence, in the former, of portions of the pancreas which are not responsible for insulin secretion.

The following is a partial reconstruction of the series of events leading to the development of metahypophyseal diabetes in the dog or in animals which react in a similar manner. It is probable that the injection of anterior pituitary extract evokes a secretion of insulin from the pancreas. Ham and Haist (16) reported an increased mitotic activity in the islet tissue of the pancreas, as well as in the thyroid parathyroid, and adrenal cortical glands following the administration of an tenor pituitary extract Weinstein (17) confirmed the earlier report of Shpiner and Soskin (18) that the injection of anterior pituitary extract may cause an immediate temporary fall in the blood sugar. The secretion of insulin in response to the antenor pituitary extract injection probably also accounts for the decreased nitrogen excretion (19, 20) However, the continuation of anterior pituitary extract treatment eventually exhausts the insulin secreting cells of the pancreas and appar ently permanently incapacitates them (10, 11) The unopposed action of the an tenor pituitary gland then becomes evident and produces an increase in protein and in fat catabolism similar to that occurring when anterior pituitary extract is injected into depancreatized animals (10)

Lukens and Dohan (o) used partially departreatized cats with metahypophyseal diabetes to study the influence of various procedures as regards their protective action on the islands of Langerhans. They found that fasting, a high fat diet. and insulin and phlorhizin administration, respectively, led to recovery from meta hypophyseal diabetes, providing the treatment were started before the insulin producing cells were completely destroyed. They pointed out that the obvious common factor in all these treatments was the maintenance of a lower blood sugar level over a period of time Best and his co-workers (21, 22, 23) had shown that fasting, high fat diets, and the administration of insulin diminished the insulin content of the pancreas of rats According to these workers, the histological pie ture of the islets of Langerhans after such treatments suggests that these pro cedures tend to put the \$-cells of the islets "at rest' Lukens and Dohan (q) adopted this suggestion to explain their own results. They concluded that in their partially depancreatized cats with limited functional reserve of the islets the administration of anterior pituitary extract led to overstimulation and exhaustion of the remaining islets through sustained hyperglycemia. The various procedures which

they employed to lower the blood sugar level presumably reduced the degree of overwork of the islets and enabled them to survive and recover

While it is difficult to offer a satisfactory alternative explanation to the above there are certain obstacles to the acceptance of the postulated mechanism. This Haist and Best reported that the insulin content of the pancreas of bypophyse tomized rats was similar to that of normal rats when both types of animal were equally well fed (23 24). If the insulin content of the pancreas were a rehable index of the rate of insulin secretion by that organ, their finding would indicate that the pancreas of the hypophysectomized animal secretes as much insulin as that of the normal animal. This would appear to be extremely unlikely, in view of the marked sensitivity of the hypophysectomized animal to administered insulin. It therefore seems hazardous to judge the state of work or rest of the pancreas on the basis of its insulin content.

As regards the influence of insulin on the histology of the islets this depends in part at least—on the experimental conditions. Mirsky (25) has shown that the continued administration of insulin to partially departeratized dogs may actually lead to the degeneration of the pancreatic remnants! Control animals with similar amounts of pancreas removed and observed for the same length of time showed no diabetes and no evidence of any developing pancreatic insufficiency. The insulin treated dogs exhibited severe acute diabetes once the insulin administration ceased and showed no tendency toward spontaneous recovery. At the present time it is not possible to reconcile these results with those of Lukens and his co workers

METATHVROID DIABETES

Houssay (26) has reported that partially depancreatized dogs given large amounts of thyroid extract over a prolonged period of time eventually exhibit a permanent diabetes similar to metahypophyseal diabetes. This substantiats the discussion in chapter sviu concerning the role of the thyroid in cabohydrate metabolism and enhances the probability that the antenor pituitary exerts its effects partly through this gland. Houssay could not produce metathyroid diabetes in dogs with the pancreas intact.

ALLOXAN DIABETES

In the course of studies on the toxic effects of alloxan (Fig. 59) on the kidney, Dunn. McLechie and Sheehan (27) noted (among other pathological landings at post mortem examination) a necrosis of the β -cells of the islands of Langerbars. Many of their ratis exhibited convulsions before death. Jacobs (38) had previously reported that the administration of alloxan caused hypoglycemia in rabbits. Dunn and co workers (27) confirmed this observation but found that some of their and roals which survived the initial effects of the drug later developed permanent diabeters.

By varying the dose of alloxan so as to avoid death from hypoglycemia and damage to tissues other than the pancreas, Bailey and Bailey (29) and Goldner and Gomori (30) were able to produce diabetes practically at will in rats, guinea pigs, rabbits, and dogs In the latter animals, kidney and hyer damage seemed to be at a minimum, the actinar tissue of the pancreas and the a-cells of the islands of Langerhans appeared to be entirely unaffected, but the β -cells of the islets were completely destroyed. These observations offer a new tool for the investigation of the diabete syndrome, particularly in small animals where complete pancrea-

ATTOYAN

Fig. 59.—Structure of gliozan, showing its close relationship to certain derivatives of naturally or curring nucleoproteins. The possibility has been suggested (33) that alloran, or a unillar substance artius from a disordered nucleoprotein metabolism may have a bearing on the ethology of disbette mellitus.

tectomy has been difficult or impossible (29, 31). It may also facilitate the study of the separate functions of the component cells of the islands of Langerhans (29, 32, 33).

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PART V

INTEGRATION OF PHYSIOLOGICAL AND CLINICAL ASPECTS



CHAPTER XYI

REGULATION OF CARBOHYDRATE METABOLISM

TE HAVE thus far dealt with the storage of carbohydrate its intercon Versions and its unixation or dissimilation by the living organism. We have seen that our knowledge of the quantitative aspects of these phe omena is rather limited. It is therefore to be expected that the development of outcus is rainer unitied. It is therefore to be expected that the development of understanding of the mechanisms which regulate carbohydrate metabolism on anouncestanding of the incuranisms which regulate caroonydrate metabonism toold be correspondingly retarded. At the present time it is impossible to predict sound be correspondingly retarded. At the present time it is impossible to predict except in the most general sort of way, what proportions of a given dose of carboeacept in the most general sort of way worst proportions of a given uose of caroo-hydrate will follow the various possible pathways for its disposal in the living or anism under a particular set of circumstances. It is impossible to calculate how action under a particular set of chemistantes at is improsume to calculate now much of the carbohydrate will be stored as glycogen, how much will be converted and how much will be dissimilated for energetic purposes

Such partitions as might be predicted are based upon empirical data from previ our partitions as inight of predicted are used upon empirical usia from previous experiments conducted under similar conditions. We know from experience one experiments conducted uniter summa conductions to a summary conductions that when a limited amount of carbohydrate is available it is likely to be used as when a timited amount or carbonyurate is available it is tikely to be used as source of energy and that little of it will appear as gly cogen or fat. It seems ob wouse of energy and that attic or a war appear as go cogen or ial. It seems on your that there must be fairly accurate mechanisms for diverting the carbohy the channel most useful for the animal but we know little or nothing of the details of such mechanisms

The regulation of the blood augar level differs somewhat from that of other and regulation of the blood augar level olders somewhat from tolat of other carbohydrate functions. Storage interconversions and dissumilation value with carbohydrate supply whereas the blood sugar level in the normal animal remains monayouate supply whereas the unous sugar reversal the monatar annual tensation of feeding and fasting. On the other hand the hypergly cenna and the great dependence of the blood sugar the date of the dabetic organism on the kind and amount of ingested food indicates a profound disturbance of the regulating mechanisms in diabetee Claude Bernard was keenly aware of the dynamic balance involved in blood

Sign regulation—the balance upon which any proper conception of regulation must be based. He clearly stated that the normal blood sugar level represented a precise equilibrium between the rates of sugar formation in the laver and of sugar invence equindrium between the rates or sugar formation in the aver aim or sugar whilst aton in the tissues (1) While the role which he assigned to the Liver has been confirmed by most recent workers (1 3 4 5 6 7 8 9) at has nevertheless been with the supplemental or clinical variables are supplemental or clinical variables are supplemental or clinical variables. states which are characterized by a persistence of abnormal blood sugar levels states which are characterized by a persistence or authorities unless sugar never instead attention has been focused almost exclusively upon the utilization of

sugar This may be accounted for partly by the discovery of insulin and partly by the erstwhile predominance of the non utilization theory of diabetes. The discovery of insulin led to overemphasis of the possible role of the pancreas in the regulation of carbohydrate metabolism, the non utilization theory demanded that the regulating activity of the pancreas be exerted upon sugar utilization.

A striking example of the manner in which these factors have influenced inter pretations is contained in a relatively recent review, in which an older paper by Pollak (10) is cited. The latter author, by fortunate deduction from meager evidence, had arrived at the conception that the blood sugar level was a determining factor as regards the activity of the liner in the regulation of carbohydrate metabolism (10, 11). The quotation from Con (12) is as follows "Pollak, before the insulinera, advocated the view that the blood sugar level is of major importance in the regulation of carbohydrate metabolism which, translated into out present ter minology, means in the secretion of insulin." It will be seen, from the evidence to be reviewed, that this "translation" is not warranted and that Pollak's version happened to be more correct.

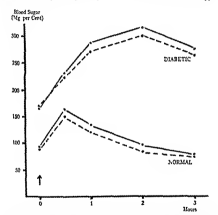
THE HOMEOSTATIC MECHANISM IN THE LIVER

The characteristic rise and fall of the blood sugar following the administration of dextrose to normal animals represents a rapid and reproducible test of the regulating mechanisms. Wide clinical and experimental use of this test has been made. In man it has been customary to have the subject drink 300-500 cc of lemon ade sweetened with 50-100 gm of dextrose. The test is usually performed in the morning before breakfast, for it has been found that previous food intake influences the outcome of the test. A control blood sugar determination is made before the test, and further determinations are made at various intervals up to 3 hours after the test. The average, or "normal," blood sugar curve obtained in the healthy subject is shown in Figure 60 where it is contrasted with the so called "diabetic" curve from patients with diabetes mellitus and from individuals suffer ing from other conditions which interfere with efficient regulation.

Until a few years ago, it was customary to explain the normal devirose tolerance curve as resulting from a stimulation of the pancreas by the administered sugar. The consequent secretion of insulin was supposed to dispose of the incoming sugar by increasing the rates of storage and "oxidation" of carbohydrates (12, 13). The abnormal type of curve characteristic of the depancreatized animal and of the diabetic human was attributed to a lack of pancreatic response, with a consequent inability to dispose of the incoming sugar at the normal rate (12, 13). It will be noted that this explanation ignored the important role of the liver in supplying sugar and the possibility that regulation might also be accomplished by controlling this supply

More recently, Soskin and his co workers (14) tested the fundamental basis of

these explanations by substituting a constant injection pump for the pancreas as the source of insulin in dogs. Completely depancreatized dogs received constant intravenous injections of insulin at rates just sufficient to maintain a normal constant blood sugar level in each particular animal. They were therefore restored to normal in a restricted experimental sense except that they could not inobiaize additional insulin, they had to get along on the constant anounts of insulin supplied



Fir 60—Oral destrose tolerance curves in normal and datebra: human. The arrow indicates the administration of 50 gm of glocose by month. The continuous lines represent arterial (capillary) blood sugar values: the rockes lines represent versous blood sugar values. Firm the data of Cavett and Schecko [17].

by their artificial substitute for a panciess. If the previous concepts had been correct, such animals should have yielded 'diabeties' dextrose tolerance curves. But, as a matter of fact, the animals exhibited perfectly normal tolerance curves. It was vident that, provided sufficient insulin seere pretent to maintain a constant blood stops let in oaddutional secretion was necessary for adequate regulation.

These results naturally directed attention toward the liver as possibly the factor that varied in regulation Normal dogs were hepatectomized, and a constant mec tion of dextrose just sufficient to maintain a normal, constant blood sugar level was substituted for the liver Since the pancreas was intact, this type of animal preparation was able to mobilize insulin as required but could not alter the rate at which sugar was being delivered to the blood from the artificial liver Such animals invariably yielded markedly "diabetic" tolerance curves. It was apparent that the pancreas was not essential to the regulating mechanisms responsible for the nor mal dextrose tolerance curve, while the presence of the normal liver was essential

This led to observations on the simultaneous blood sugar values of the blood flowing into and out of the liver, in normal and departreatized dogs, during the course of dextrose tolerance tests. From these and the previous results it was postulated that (in the presence of a sufficiency of insulin, but not necessarily an extra secretion from the pancreas) the normal liver, as one of its responses to ad ministered dextrose, decreases the output of blood sugar which it has previously been supplying from its own resources

The homeostatic regulating mechanism for the control of the blood sugar level was later subjected to direct proof (15) By correlating the rate of blood flow through the liver of experimental animals with the difference in the sugar content between the blood flowing into and out of this organ, it was possible to calculate the absolute amounts of sugar entering and leaving the liver per unit of time Fig ure 61 illustrates such an experiment and shows what happens when a dextrose tolerance test is made. It may be seen that the liver, which was pouring sugar into the blood prior to the administration of the dextrose, ceased to do so almost im mediately upon the administration of dextrose and started to take in large quan tities of sugar (The period following this retention of sugar is particularly worthy of note At this time the liver neither took in nor put out sugar for a period of about an hour, showing that the inhibition of the output of sugar is a phenomenon separate from the storage of sugar) When the period of inhibition was over, the liver again began its usual supply of sugar to the blood, and the blood sugar level which had fallen somewhat below the pre test level during the inhibition rose up to and slightly above its pre-test level

In further experiments (16) it was also shown that completely deparcreatized dogs which were receiving the appropriate constant injections of insulin exhibited at least as great a hypoglycemic reaction following the cessation of prolonged sugar administration as did normal dogs (see Fig 62) Like the normal dextrose toler ance curve, this phenomenon cannot be ascribed to insulin mobilization but must be accounted for by the decrease in the output of sugar by the liver in response to the influx of exogenous sugar In other words, this period of bypoglycemia fol lowing the dextrose tolerance curve or following the cessation of more prolonged dextrose injections corresponds to the time which elapses before the liver is able

to accelerate its rate of supply of blood sugar to a point sufficient to maintain the onginal normal blood sugar level 251

rignau normai nuouu sugar ievei The hepatic regulating mechanism is analogous to the system used for the regu THE REPORTER REQUIREMENT AND AMERICAN USES THE STREET ASSETTING THE REPORT OF THE PROPERTY OF sation or temperature in many modern mones, manney, the intermostat numate ar rangement. When the temperature of the house rises above the level at which the rangement. When the temperature on the nouse also above the reverse which the themsets that been set the furnace shuts off until the excess heat has been dissipated by the control of the pated When the temperature of the house falls back to the threshold of the ther mostat, the furnace starts up again. That is exactly what the liver does, so far as (Mg per Cent) (Mg per Min)

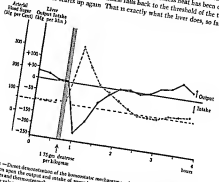


Fig. 64.—Direct demonstration of the homeostatic mechanism in the liver. The effect of destroyed and state has been the large of a part of the property of the large of a part of the property of the large of the la Fo de - Direct demonstration of the homeostate mechanism in the laver. The effect of destroyed ministration specifies and state of signs by the laver of an intext dog; activities and shared states of the state of constitution upon the output and intake of sugar by the laves of an intake dog calculated from blood aggressive, and thermostromoute measurements of departs blood don. The broken has represent a created and aggressive and the process of the proce Sign values and thermostromular measurement of departs blood flow. The broken has represent actions for any or state of sugar by the area of the sugar cost money have present output or state of sugar by the area continued to the sugar by the area of sugar by th soon user values the heavyer cont mous hier represents output or matate or sugar by the later in multi-fring per minute. Vote the numerical cressions of sugar output when sugar as distincted and her later angles of sugar and the second base after wave administration by the later in multifrom per panule. Vote the immediate cessation of sagar output when sagar is administered and the single indicate of ungar which follows. Throughout the second hour after sugar administration the lines of the sagar in the sternal should fail the sagar in the sternal should fail be low in kee intake of ugar which follows. Throughout the second hour after sugar administration the liver outlier retains nor excrete sugar. During this proof the level of sugar in the arternal blood falls below our many control with the second control after the level has returned to solve the terms. Softer relates nor exercise sour. During this period the level of sugar in the arterial bised falls below its common control values and does not return to atomat until after the her has returned to other Designation of the housest control of the common of the ontail one of values and does not return to sormal until after the aver has removed its output. The in of super is therefore a real and separate phenomenous from the storage. of sugar (Soskin et al [15])

the blood sugar level is concerned in this analogy the temperature is equivalent to the blood sugar level and the thermostat furnace arrangement is represented by the her It will be noted that, just as it is the room temperature which operates og menter at with be noted that, just as it is the foom temperature which operate the thermostat and shuts off the furnace so it is the blood sugar level which in hibits the output of sugar by the liver

Accordingly, the dextrose tolerance curve and the hypoglycemic phase which often follows it resemble the fluctuations in temperature above and below the threshold of regulation when an extra quantity of heat is introduced into the tem perature regulated house. The characteristics of the curve depend upon the mag intude of the disturbing factor (the amount of sugar administered), the setting and sensitivity of the thermostat (the endocrine halance), and the capacity of the furnace (the ability of the liver to produce sugar).

The fact that the hepatectomized animal with an artificially maintained normal constant blood sugar level (and with the pancreas and extrahepatic tissues free to

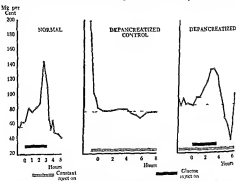


Fig. 62—Hypoglycemic reaction without extra insular. The solid black line labeled. Gloscoe layer tion refers to the injection of the test sugar. The crosslatched line labeled. Constant injection of insular place serious constant injection of insular place serious that are required to maintain a normal blood signal red.

exert whatever regulating powers they possess) yields 'diabetic dextrose toler ance curves (14)' indicates the essential role of the liver in blood sugar regulation

It is not to be supposed, however, that the hepatic mechanism is the only one involved Glycogen deposition in both the liver and muscle and an increased utilization of sugar by the extrahepatic tissues undoubtedly play their parts. These processes, like hepatic homeostasis, are under the influence of the blood sugar level Cori and Cori (17) have pointed out that the rate of glycogen deposition depends upon the concentration of sugar in the blood Soshim and Levine (18) have shown that the rate of sugar utilization by the extrahepatic tissues varies directly with the height of the blood sugar level It seems blorgal to assume that smaller amounts of sugar, especially if they enter the circulation via the portal vein, may be fully compensated for by hepatic lighbition alone. Larger amounts of sugar will invoke hepatic storage as well Still Jareer amounts, which, in soite of the forecome, raise

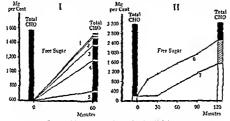


Fig. 63—Inhibition of liver glycogenolysis by added glucose

I The industrie of different amounts of glucose added to each vessel upon the appearance of free larger ni area.

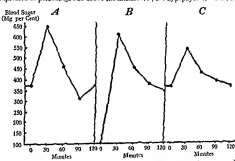
The blocks representing total carbohydrate determinations at the beginning and end of each experiment and cate that there was no significant loss of exthohydrate from the system (Soskin et al. [10])

the systemic blood sugar level, will bring into play the additional factors of extrahepatic storage and increased utilization

It is clear that the fundamental regulation of the blood sugar is an autoregula too, in which the prime mover is the blood sugar level itself. This is further supported by the movin of Soskin, Levine, and Taubenhaus (19) on the rate of appearance of free sugar in glycogenolyzing liver bere with and without the presence of added destrose. The results are illustrated by Figure 6; It may be seen that the sugar level influences the enzyme system concerned with the Glycogen=Glucose

(23) rather than an index of the ability to handle sugar, once it has entered the blood stream

However, even when the sugar is administered intravenously, the apparent tol crance depends upon how the data are expressed. Figure 65 shows a typical intravenous dextrose tolerance curve in a hypophysectomized dog (broken line) as compared to a similar test in a normal dog (continuous line). If the lower initial and final levels in the hypophysectomized animal are ignored, its curve appears to be low or better than normal. If, on the other hand, the results of both curves are expressed as percentage rise above the initial level, the hypophysectomized curve.

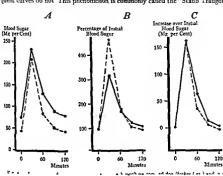


Fir 64 -- Normal dextrose tolerance curves in the 'Houssay' dog Dextrose tolerance curves ob

appears to be high or worse than normal. When, however, the actual curves are drawn from the same base line it can be seen that they are practically identical

meterica.

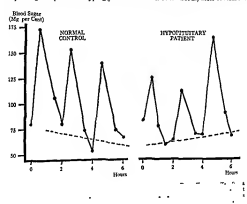
The "triple tolerance test"—Although the principal regulating action of the liver snormal in the hypophysectomized animal there is a subsidiary regulating mech insim which is not normal, namely, that mechanism which is due to the presence of the pituitary itself. When dextrose tolerance tests are repeated in a normal anial, each test starting as soon as the previous one is over, it will be found that the second curve is lower or better than the first, while the third is usually better than the second. The fourth curve may show some further improvement, but subsequent curves do not. This phenomenon is commonly called the "Staub Traupott



effect," after the investigators who first described it (24). It has been shown that this phenomenon does not occur in the hypophysectomized animal (25).

In the absence of the hypophysis the first curve is the lowest or best one ob

tained (Fig 66) It might be supposed that this abnormality is due to some second ary effect of the absence of anterior pituitary secretion upon the function of the liver. But this is not the case, for the administration of anterior pituitary extract to the hypophysectomized animal raises the level of all the tests without restoring the Staub Traugott phenomenon. The lowering of the second and third curves (and sometimes the fourth) in the normal animal can therefore best be explained as a progressive depression in the activity of the pituitary gland, as a result of repeated or prolonged exposure to hyperglycemic levels. In other words, after several suc



cessive doses of sugar the normal animal reaches that stage at which the hypoph ysectomized animal starts out. This mechanism is very acceptable from the felo logic standpoint for it is obvious that, during continued high sugar intake, regulation will be more efficient as the threshold of the mechanism is lowered. It is equivalent to the common practice of setting down the thermostat of the house to say 50°F during the spring or fall months, when only an occasional, brief cold snap may be expected.

Influence of the adrenal cortex and the thyroid gland —At the present time it is injuence of the advenas cover and the sayrous grand—At the present time it is difficult to separate the influences of the adrenal cortex and the thyroid gland from amount to separate the innuences of the aurenal cortex and the entyroid giand from that of the anterior pituitary. Indeed, some of the influence of the anterior pituit 250 that of the anterior pituitary indeed some of the innuence of the anterior pituitary gland described above may be exerted through these other glands (26 27) say ganu uescrineu anove may ne exerteu tinougu these other grants (20 27). At any rate deficiency or removal of the adrenal cortex on the one hand or the At any rate denotency or removal of the advenue cortex on the one nano or the administration of potent extracts of this gland on the other hand will lower or auministration or potent extracts of this grand on the other hand will lower or raise the blood sugar level in a manner resembling that which occurs when the raise toe otooo sugar rever in a manner resembling that which occurs when the pituitary hormone is varied. To a lesser extent, this is also true of the thy roid (28). productly authors to value a conserver extent this is about the or the mytom (20). Presumably, then, the adrenal cortex and the thyroid influence the threshold of regulation of the sugar level in the same manner as does the antenor pituitary

influence of the state of the liver on the regulation of

Although we have compared the liver to a thermostat furnace arrangement we numous we have compared the aver to a thermostal furnace arrangement we have thus far considered only those factors which operate by affecting the thermostal furnace arrangement we more units let considered only those factors which operate by allecting the intermo-tial part of the mechanism. However, it is obvious that regardless of where the seat part of the mechanism thowever it is obvious that regardless of where the ihemostat is set—the state of repair and the capabilities of the furnace will have in important bearing on the degree of regulation which is achieved. For example, on important ocating of 80° f would have no meaning if the furnace were incapable a menting at the stitung of so 1 wound have no meaning it the suinage were incapation of producing enough heat to raise the temperature of the house to that level An on producting enough near to raise the temperature of the nouse to that here can other consideration is the speed with which the rate of heat production by the furnace can be increased or diminished. Unless such adjustments are rapid there will nace van de increased or diminished. Unless such adjustments are rapid. Incre will be a considerable overswing before the correct temperature is reached. If the ther mostat on a sluggish furnace checks over at let us say 80°F the temperature may meets of or 100° l before the effect of shutting off the furnace becomes evident Finally, even with a furnace of great capacity and high efficiency, the degree of regulation will depend upon the magnitude of the environmental temperature Association will depend upon the magnitude of the environmental temperature change for which the furnace has to compensate. In other words, the usual nightly dop of 10-20°F in the outside temperature might produce practically no per copible disturbance in the temperature of the house while a sudden frost drop mpms the outside temperature 40° 50° F might result in a downward dip in the one courside temperature 40° 50° r might result in a downward dip in the house temperature before the furnace could cope with it. The analogous considera lions apply to the liver as the organ which makes the blood sugar

was apply to the liver as the organ which makes the mood sugar. An example of a disturbance in sugar regulation analogous to the situation in shirth the furnace is incapable of raising the temperature up to the level at which auce the turnace is incapable of raising the temperature up to the reversal which be thermostat is set is the effect of fasting on the hypophysectomized animal and on the hypopututary human (29) The withholding of food in the latter organisms of the hypopituitary human (29). The withnoising of tool in the latter organisms trouble in a progressive hypoglycemia. This does not depend upon any change in regulation because the resumption of food intake immediately restores the previous ous blood sugar level. It does depend upon a marked reduction in the ability of the liver to make blood sugar from body stores so that it cannot supply sufficient

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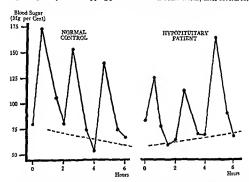


Fig. 65 — Consecutive destrow tolerance curves at a hour intervals (*Triple Tolerance Text.) in a normal human and in a proved case of hypopotimism. Therety five agrains of destroise in specific aqueous solution was injected intravenously in every instance. Note the sharp contrast between the slopes of the lowest points in each sterse as understed by the bubbers hum? This text has been found to be a use full objective criterion in the study of proved and suspected cases of hypopituitarism in humans a bea used in conjunction with clinic all data. (Sokim [53])

cessive doses of sugar the normal animal reaches that stage at which the hypoph ysectomized animal starts out. This mechanism is very acceptable from the teleo logic standpoint, for it is obvious that, during continued high sugar intake, regulation will be more efficient as the threshold of the mechanism is lowered. It is equivalent to the common practice of setting down the thermostat of the house to, say, 50° F during the spring or fall mouths when only an occasional, brief cold snap may be expected.

Influence of the adrenal cortex and the thyroid gland —At the present time it is difficult to separate the influences of the adrenal cortex and the thyroid gland from that of the anterior pituitary. Indeed, some of the influence of the anterior pituitary gland described above may be cretted through these other glands (26, 27). At any rate, deficiency or removal of the adrenal cortex, on the one hand, or the administration of potent extracts of this gland, on the other hand, will lower or raise the blood sugar level in a manner resembling that which occurs when the pituitary hormone is varied. To a lesser extent, this is also true of the thyroid (26). Presumably, then, the adrenal cortex and the thyroid influence the threshold of regulation of the sugar level in the same manner as does the anterior pituitary.

INFLUENCE OF THE STATE OF THE LIVER ON THE REGULATION OF THE BLOOD SUGAR

Although we have compared the liver to a thermostat furnace arrangement, we bave thus far considered only those factors which operate by affecting the thermostat part of the mechanism However, it is obvious that, regardless of where the thermostat is set, the state of repair and the capabilities of the furnace will have an important bearing on the degree of regulation which is achieved. For example, a thermostat setting of 80° F would have no meaning if the furnace were incapable of producing enough heat to raise the temperature of the house to that level Another consideration is the speed with which the rate of heat production by the furnace can be increased or diminished. Unless such adjustments are rapid, there will be a considerable overswing before the correct temperature is reached. If the thermostat on a sluggish furnace clicks over at, let us say, 80° F, the temperature may rise to go or 100° I before the effect of shutting off the furnace becomes evident Finally, even with a furnace of great capacity and high efficiency, the degree of regulation will depend upon the magnitude of the environmental temperature change for which the furnace has to compensate In other words, the usual nightly drop of 10°-20° F in the outside temperature might produce practically no per ceptible disturbance in the temperature of the house, while a sudden frost, dropping the outside temperature 40°-50° F, might result in a downward dip in the house temperature before the furnace could cope with it. The analogous considera tions apply to the liver as the organ which makes the blood sugar

An example of a disturbance in sugar regulation analogous to the situation in which the furnace is incapable of raising the temperature up to the level at which the thermost is set is the effect of fasting on the hypophysectomized animal and on the hypopituitary human (20). The withholding of food in the latter organisms results in a progressive hypogly cenia. This does not depend upon any change in regulation, because the resumption of food intake immediately restores the previous blood sugar level. It does depend upon any cannot supply sufficient here to make blood sugar from body stores, so that it cannot supply sufficient

sugar to maintain the blood sugar level unless additional preformed sugar or amino acids regularly enter from the gastro intestinal tract (27)

The situation in the liver which is analogous to the sluggish furnace, unable to increase or decrease its rates of heat production very readily, is that where the liver is damaged by toxic agents It is well known that the "diabetic' type of dex trose tolerance curve is obtained in this condition (so at)

The "daabette' type of tolerance curve obtained in starvation or on a high lat diet is analogous to the temporary breakdown in the temperature regulation of the house when a sudden great demand is made upon even a very efficient furnace. Both starvation and fat feeding are alike in that no preformed carbohydrate is being received by the body, so that the liver must make all the necessary carbohydrate from its own resources. This represents a high degree of activity on the part of the liver, as compared to the normal conditions, under which it need manu facture only a small proportion of the body's requirements. The deceleration of sugar output by the liver when sugar is administered requires a longer time when the liver is working at top speed than when it is working at half or quarter speed. The essential correctness of this interpretation is supported by the fact that it is only the first does of sugar given to a starved or fat fed animal that results in the "diabetic type of curve. The second dose (by which time the liver has been able to slow up its production) usually shows a return of the dextrose tolerance curve toward the normal (12).

ACTUAL COMPLEXITY OF REGULATION IN THE LIVING ORGANISM

Thus far, the analogy of the thermostat furnace arrangement has served us well in helping to simplify the relationship between the endocrine glands and the liver in the regulation of the blood sugar. But it is necessary to realize that the mechanism which has been described is integrated with a series of other regulatory processes in the body. We have said, for example, that the threshold of regulation of the liver is determined by the endocrine balance. But what determines the character stite rates of activity of the endocrine glands which maintain this balance? This

that the blood sugar level affects not only the liver but also the activity of the letter printitary gland which in turn influences the reaction of the liver to the blood sugar level (22). There is also evidence that the concentration of sugar in the blood passing through the pancreas influences the rate of secretion of insulin (33). Fur thermore, the concentration of a given hormone in the blood may have a control ling action upon the activity of the gland which secretes that hormone (34). An other mode of regulation may occur by the controlling effect of the hormone of one gland upon the rate of activity of another gland. An example of the latter type of

effect is the excessive stimulation of the secretion of insulin by the repeated intection of massive doses of extracts of the anterior pituitary gland exentual) ? . to islet exhaustion and pancreatic diabetes as firet

It is not unusual in the .

oing mechanisms all dur

capable of serving the

function to a considerah at the other mechanisms are impaired by dis ease or by an experimental procedure. This situation exists in recard to the regula tion of the blood sugar. It has been possible to demonstrate a primitive type of regulation of sugar output by the liver which can occur in isolated hepatic tissue in the test tube (19) (see p 253) In other words the output of sugar is to a certain extent controlled by the concentration of sugar present even in the absence of any possible endocrine adjustment. In addition to this intrinsic hepatic mechan ism and its endocrine regulators which have already been discussed there are also certain emergency mechanisms mediated by the central nervous system and the adrenal medulla (see chan xv p 168) The latter mechanisms are not evident under normal conditions and they can be entirely eliminated experimentally without appreciably affecting the sensitivity of regulation. But when under ah normal conditions of stress and strain the organism is threatened by an unduly rap d or profound hypoglycemia the emergency mechanisms rapidly come into play by breaking down liver gly cogen and providing the needed blood sugar

It may be helpful to think of the relationships between the emergency mecha n sms the endocrine glands and the intrinsic hepatic homeostas s from the phylo genetic viewpoint. The fundamental or primitive regulation may be supposed to reside in the brochemical processes of the tissue cells. The endocrine glands may represent a step up the evolutionary scale by providing a more sensitive and finely adjusted regulating mechanism, which renders the more highly developed organ ism less dependent upon its external environment. The emergency mechanisms may be an add tional protection against hypoglycemia for the highly specialized tissues (e.g. central nervous system) of the most highly developed organisms

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CHAPTER XXII

PATHOLOGICAL PHYSIOLOGY AND CLINICAL APPLICATIONS

FIER having outlined the influences of the various endocrine glands up the process of blood sugar regulation which occurs primarily in the liv It becomes a relatively simple matter to account for the characters chinical disturbances which accompany disease or dysfunction of the glands or the liver

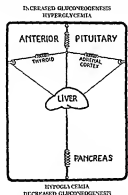
CLINICAL DISTURBANCES IN THE ENDOCRINE REGULATION OF THE BLOOD SUGAR

We have seen that the experimental diabetic syndrome is primarily a d turbance in the regulation of carbohydrate metabolism (rather than of utilization brought about by various manipulations of the endocrine glands or their hormone But in order to avoid confusion in terminology, it is necessary to remember at the outset that diabetes mellitus, as it occurs in man, is still a clinical syndrome of u known etiology. The essential and minimal characteristics of this syndrome are persistent hypergly cerms with gly cosums-all other effects, such as polyuna, d hydration, demineralization, loss of weight, ketosis, and coma being secondary (1 In the mildest disturbances the diagnosis of diabetes mellitus often cannot b finally established until the condition has progressed in severity to the point that stable persistent criteria develop. It often happens, also, that a mild disturbance in carbohydrate regulation is found to be accompanied by hepatic damage, hyper thyroidism, adrenal cortical tumor etc. If the liver disease or the glandular dis turbance is adequately treated by medical or surgical means and the carbohydrat disturbance is thereby eliminated, it is not customary to label the transitory by perglycemia and glycosuma as diabetes mellitus

It is readily understood that the foregoing terminology is merely a chinical con vention. From the physiologic standpoint it is difficult to conceive of a disturbance like diabetes mellitus, which, in some individuals, would not be found in minimal and transitory form. Nor does the presence of frank and remediable liver disease or glandular disturbance necessarily make the resulting diabetes any different from that which occurs when the etiologic disturbance cannot be detected by present chinical methods. It is this physiologic point of view which must be kept in mind in considering the possible etiologic factors involved in the recognized clin-

ical disturbance

Since the condition, which by clinical convention is called "diabetes mellitus," is characterized, at the present time, by the very lack of any consistent demonstra ble abnormality in the endocrine glands," we must perforce base our notions as to possible etiology upon the various experimental procedures by which a similar syndrome can be produced. These possibilities have already been indicated in the extinois devoted to the various endocrine glands and the liver. Their relationships to each other are graphically illustrated in Figure 67. In the balance of forces rep-



DECREASED GLUCONEOGENESIS

Fto 67 —Mechanical analogy to the endocrine balance

resented there, it may readily be seen that the same end result might be obtained in a variety of ways. A shift of regulation toward hypergij cemia might be due to a diminution in the insulin factor (an absolute lack of insulin) or to an intensifica

Two recent publications require some comment

¹ Susman (108) has reported camera lucida measurements of the relative areas of the islets of Langer hans in histologic sections of panerestic glands from human belogs with and without dabetes meditus. According to him the islets of the diabetic individuals accopied of 17-4 6 per cent of the total area, as

tion of the opposing factors (a relative lack of insulin) If the latter type of dis turbance is, indeed, responsible for some cases of diabetes mellitus, it is possible that we may eventually learn to distinguish a pituitary diabetes, an adrenal corti cal diabetes and a thyroid diabetes, as well as a pancreatic diabetes. To this list must be added a possible hepatic diabetes which might occur in the absence of endocrine disturbance when the liver is no longer responding normally to its endocrine regulation It must be emphasized that none of these considerations minimizes the importance of insulin in therapy or suggests that any other efficacious agent is known at the present time The diagram clearly indicates that the important thing from the therapeutic standpoint, is the maintenance of the normal balance The administration of insulin will correct the imbalance whether it is due to an absolute or to a relative lack of this hormone

The differentiation of the various possible types of diabetes mellitus must await the development of adequate methods for the quantitative estimation of glandular function or of the titer of the various hormones in the blood. For the present all diabetic manifestations which are accompanied by a clinically recognizable dys function of some gland or of the liver are considered to he part of the syndrome as sociated with that clinical state. A similar situation exists as regards carbohydrate disturbances in the direction of hypoglycemia and the differentiation between hypermsulmism and other conditions which may lead to hypoglycemia. An inspec tion of the following list, in conjunction with an examination of Figure 67 will re late the characteristic blood sugar disturbances accompanying the various known endocrine syndromes with the physiologic considerations which have been out lined. We have included key references to articles dealing with the carbohydrate disturbance in the clinical syndrome

EMPOCRANE HYPERGLYCENIAS

Anterior pituitary

Acromegaly (2)

Pituitary basophilism (3 4) Thyroid Hyperthyroidism (5)

Adrenal cortex Hyperadrenocorticalism (6 7)

Pheochromocytoma (8)

Adrenal medulia

Diabetes mellitus in those cases where there is evi Pancreas dence of destruction of the islets of Langerhans (a)

compared to 0.7.5 sper cent in the pastress of normal individual. Aside from the cons detable overlap in these figures it should be pointed out that there is a difference of only 56 per cent in the mean value of the constraints. The constraints of the constrai

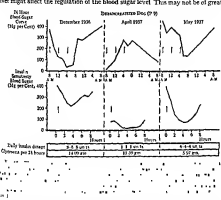
ENDOCRINE HYPOGLYCENIAS

Anterior pituitary Summonds' disease (10)
Anorexia nervosa (11)

Thyroid Hypothyroidism (12)
Adrenal cortex
Addison a disease (13)
Adrenal apoplexy (14)
Pancreas
Hyperinsulnism (14)

INFLUENCE OF LIVER DYSFUNCTION ON BLOOD SUGAR REGULATION

In chapter xxx (p 259) we described the various ways in which the state of the liver might affect the regulation of the blood sugar level. This may not be of great



practical importance when one is dealing clinically with a case of frank liver disrese, where the danger to the from other consequences of liver failure overshadows the carboby drate disturbance. But it may be of considerable value in diagnosis and prognosis when an endocrine disturbance in blood sugar regulation is compilcated by the presence of liver dysfunction. Figure 68 illustrates a striking example of this situation. Here we have a pure endocrine disorder, namely, diabetes re sulting from the removal of the pancreas in the dog, experimentally complicated by a reversible type of liver damage (16) It will be seen that the characteristics of the diabetes in this dog were markedly changed during the time that the liver was affected (April, 1937)

Interest in these results is enhanced by the fact that in chincal diabetes mellitus we find two similar types of the disease—namely, the insulin sensitive ("adult" or stable). The depancreatized dog with an unimpaired livet (December, 1936) resembles the individual with insulin sensitive, juvenile, or instable diabetes mellitus. The morning fasting blood sugar is the highest in the 24 hours, the blood sugar list sharply during the day under the influence of a dose of insulin with each meal and then rises throughout the might hours. In this state the administration of o 3 units of in sulin per kilogram of body weight causes a smart fall in the blood sugar level of about 200 mp per cent.

The same animal, which had been on a diet of lean meat, sugar, and raw pan creas, was then placed on an equicalone due from which the pancreas was omitted. This is known to result in a severe faith infiltration of the liver [17, 18] (see chap viui, p q1]. The impairment of liver function consequent to the faith infiltration is reflected in three ways which are characteristic of the insulin insensitive adult.

of the food intake, despite the insulin administered with the meals 1 ne work sugar then falls during the might hours. The administration of the same amount of mistlin as in the previous senaturity test now results in a much smaller drop in the blood sugar level. The restoration of raw paneters to the diet of this animal, with a return of the liver function almost to normal (May, 1937), completely reverses the nature of the diabetes to its original condition.

This demonstration of the influence of fatty infiltration of the liver on the nature and severity of diabetic manifestations suggests an explanation for the partial success of the extreme high fat diets and starvation regumen formedly used in the treatment of diabetes mellitus. Both these procedures will lead to a fatty infiltration of the liver. It should be noted, however, that the diabetes is controlled only at the expense of liver function. Hence it may be said that "the diabetes is testre but the patient is worse. The lack of general well being of patients under those treatments, as compared to patients under modern treatment, may well be ascribed to the difference in the functional state of the liver (120).

Toxemic liner damage—Abnormal dextrose tolerance curves have been de scribed as occurring in patients suffering from acute infectious diseases (20) Similar disturbances in carbohydrate metabohim bave been demonstrated in expect mentally induced toxemias in animals (21, 22) The "diabetic" type of dextrose

tolerance curve obtained under these circumstances has been interpreted by some as being due to a lack of endogenous insulin, consequent to the functional impairment of the islands of Langerhams (20) Others have ascribed the phenomenon to an interference with the action of the available insulin, whether of endogenous or of exceenous orient (23)

Using methods similar to those which they employed in demonstrating the homosticum mechanism for blood sugar regulation (chap xm, p 348), Soskin and his co-workers (24) showed that toxemia affects ear-holydrate metabolism hy dam sang the liver and interfering with its regulating mechanism. Completely deparametrized dogs receiving a constant injection of insulin sufficient to maintain a correctated dogs receiving a constant injection of insulin sufficient to maintain a corrective of the constant injection of insulin sufficient to maintain a corrective of the constant injection of insulin sufficient to maintain a corrective of the constant injection of insulin sufficient to maintain a correction of the constant injection of insulin sufficient to maintain a correction of the constant injection of the constant inject

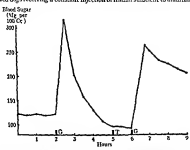


Fig. 69— D abetic. Offerance curve resulting from toom as absence of parcrias. The dog recurved from the represent a constant spection of determing the sushing pair sufficient to minima the blood signs at a constant level. G and cates the administration of the test signs and T the administration of the test signs are significant to the sign are sign are

stant normal blood sugar level were rendered toxemic by the intra-enous adminitration of diphtheria toxin. Figure 69 shows that such animals exhibit normal dextrose-tolerance curves before, and "diabetic" curve safter, town administration. Hence the abnormal tolerance curves eannot be ascribed to an effect of the toxin on the pancreas. There is also direct in vitre evidence of the influence of toxins on carbobydrate metabolism in the liver (25).

Although the "diabetic" type of dextrose tolerance curve is usually obtained in toxeme states, Althousen and others (a6) have shown that in less acute toxemiss, where there was a longer survival period, the "diabetic" type of curve may give way to the "supernormal" before death intervenes Clinically this variation in the

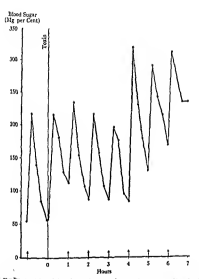
abnormal curve caused by liver damage has been described by Judd et al. (27), and it is well known that "diabetic," "supernormal," and even "normal dextrise tol erance curves may be obtained in cases of liver injury without apparent relation to the degree of liver damage as judged by clinical or pathologic criteria. Indeed this lack of correlation has been reported by Mann (28) as also applying to other tests of liver function. However the foregoing variations in response are not as haphazard as they appear but depend upon the stage or degree of liver damage which exists at the time the test is neuformed.

When a slowly progressive toxemia is induced in experimental animals and tol erance curves are repeated consecutively to the point of death (29), a definite and predictable sequence of tests is obtained, as shown in Figure 70 The first effect of the toxin is to cause a "diabetic" type of curve As the toxemia progresses there is a reversal of effect, so that the curves appear to be more and more "normal" As death approaches, there is a sudden change back to the "diabetic' type of re sponse The sequence of events portrayed in Figure 70 was obtained when 0 9 gm of dextrose per kilogram of body weight, administered intratenously, was used as the test dose of sugar The significance of the responses becomes apparent only when they are compared with those obtained using smaller and larger test doses When this is done, it becomes evident that the "diabetic" curves obtained in early toxemia are due to an impairment of the responsiveness of the bepatic homeostatic mechanism, for at this stage a small test dose of sugar (o 25 gm/kg) yields an earli er and more "diabetic" response than a large test dose (1 75 gm/kg) On the other hand, the "diabetic' type of curve obtained in late toxemia has little relationship to the homeostatic mechanism but may rather be ascribed to advanced liver fall ure At this stage the animal responds to the dextrose tolerance test in a manner similar to that of the hepatectomized animal (chap xxi, p 252) The small test dose of sugar yields normal appearing curves, while the larger test doses give pro gressively more "diabetic" curves

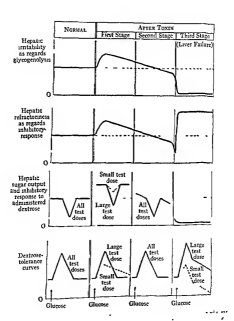
Figure 71 diagrammatically summarizes the progressive change in liver response to administered sugar. This may be explained on the basis that the first effect of a poison on the liver is to act as an irritant to the glycogenolytic mecha

> ards iven

rise in blood sugar would inhibit this process. As the effects of the toxin on the liver progress to the point of mortal damage to the hepatic cells, the latter must pass from the stage of glycogenolytic byperintability, through normal intability, to hypo-intability and death. Translated into terms of the inhibitory reaction which determines the character of the dextrose tolerance curve, this cycle of events would be (i) a decreased inhibition of glycogenolysis yielding "diabetic" deterance curves, unless the strength of the stimulus, as represented by the admin



No 70 ~ Progressive tozemic liver damage. Successive destrole tolerance curves obtained with 0.9 m. of destrole per historium of body weight administred metwoodly. Initial control curve is followed y tom administration. Arrows represent sugar administration. (Soskin and Mirsky [26].)



istered sugar, be great enough to overcome the refractory state of the organ, when a normal inhibitory response and therefore a normal tolerance curve may be obtained, (2) a return to the normal inhibitory reaction yielding apparently normal curves, and (3) a transitory phase of increased inhibition of gly cogenolysis yield may supernormal curves, which passes rapidly into the stage of complete fine fail ure, with a cessation of sugar output and the reactions of the hepatectomized am mat.

From the practical standpoint it is noteworthy that a supposedly normal dex toss-tolerance curve may under appropriate circumstances represent a greater degree of liver damage than a 'diabetic' curve. This probably accounts for the difficulty in correlating results of dextross-tolerance tests with the clinical or pathological evidence of liver damage. Such curves can be more correctly interpreted in the light of the cycle of events described above and in conjunction with other evidence as to the ertent and duration of the bepatic impairment.

It is evident that a sense of dextrose tolerance tests performed at intervals during the course of a hepatic disorder can juick information of greater prognosis value than could possibly be derived from any single test. It is also likely that a companison of tolerance curies obtained with large and small doses of sugar might be of clinical value, since in stage 1 the large dose yields more normal curves than doses the small dose, while in stage 3 the reverse is true. In general stage 1 corresponds to the carbohy drate abnormalities observed in so-called hepatitis (31, 31), while the disturbances described for stage 3 are seen in advanced hepatic curboss (33, 44).

Holmes (35) has reviewed the 18 tile observations upon the effects of toxin on carbohydrate metabolism of liver. The results of such work confirm the experimental and climical observations detailed above. The progressive effects demon strated on liver slices and arranged in order of time sequence or of degrees of damage, are as follows first, an increased rate of glycogenoly six and a decreased shibly to form glycogen from glocose (sugar can still be made from lactic and py nive soids and from alanine but cannot be stored) and second a decreased ability to form the ability to form glycogen from glycogen or less complete loss of the ability to form glycogen.

THE INTRAVENOUS DEXTROSE TOLERANCE TEST FOR LIVER DYSPUNCTION

The important influence of the state of the liver on blood sugar regulation makes it desirable to be able to differentiate between hepatic and endocrine disturbances. There have been a number of uncestigators who have reported that the ori dectates clearance curve is subnormal in hier disease, but no characteristics which would distinguish such a curve from that obtained in disbects mellitus have ever been described (11, 24). Using a standardured intravenous procedure for the test, Soskin and his co-portlers (ct) have recently been able to obtain curves from

normal individuals, patients with known liver disease, and patients with mild dathetes melititis, respectively, which are characteristic for each condition and which can be differentiated from each other. The procedure which must be followed exactly if their standards are to be used, is as follows. The test is done in the morning before breakfast. One third gram of dextrose per kilogram of body weight, in a 50 per cent aqueous solution, is injected intravenously within a period of 3 5 minutes. Blood samples are taken before the sugar administration and at 3, 1 and 2 hours thereafter. These investigators used capillary blood obtained by

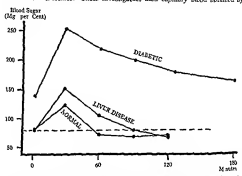


Fig. 21 — The average intravenous destrose tolerance curves of 30 hornal control individuals as of the middet nears of disbetes quellins than were available and 50 cases of proved mid of early lived case. The hornal curve returns to the pre-superior level by 60 m autes the bepatic curve returns after 50 and before 120 minutes the darbetic curve returns after 250 minutes (Socking 153).

finger puncture, and the uncromodification of the Somogyi Shaffer Hartmann method for true blood sugar

Figure 72 shows the average curves for 30 normal control individuals 25 of the muldest cases of diabetes mellitus which were available (none had a fasting blood sugar level over 200 mg per cent and none required insulin for the control of their diabetes), and 50 cases of mild or early liver disease (clinically established and corroborated by several laboratory criteria). The wide spread between the three types of curve and the ease with which they can be differentiated as apparent As regards the variation between the individual tests which go to make up the aver

age curves not a single one of the 30 normal cases took as much as 60 minutes to return to the pre injection level. This agrees with the normal standard previously reported by Tunbindge and Allibone (30). Not a single one of the 25 cases of mild dabetes took, less than 120 minutes to return to the initial level. Not a single one of the 30 cases of mild or early liver disease took as long as 120 minutes to return to the pre injection level, although 13 of the 50 or approximately 25 per cent of these cases did cross the have high no less than 60 minutes.

It might appear, at first glance from the average curves, that the differentiation between the diabetic and hepatic type can just as readily be made from the higher

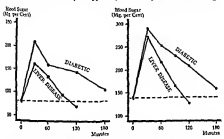


Fig. 23—Individual intervenous-destrose solerance curves in cases of diabetes and of liver disease which the provide to start at identical fasting blood sugar levels. The characteristic downshops and the limes of return to the pre injection levels are the enterior for different ation. (Sokin [13]).

initial level and the higher peal, value of the former. This is not so when individual curves are considered. The characteristics of the average curves depend upon the lact that more of the disabetic curves started at, and reached, higher level evis. However, the range of these values in disabetes and in liver disease actually overlapped to some extent. Figure 73 shows that, when this was the case, the characteristic downward slope of the curve and the time at which it crossed the base line were the real differentiating factors. The curves in Figure 73 are for individual cases of disabetes and of hier disease, selected because they happened to start at identical fisting blood sugar levels. It may be seen that, while the initial levels and highest peaks did not distinguish between the two conditions. The characteristic down slope and time of return to the pre injection level permitted easy distinction.

THERAPEUTIC USE OF HIGH CARBOHYDRATE DIETS IN LIVER DISEASE

Since Rosenbaum (37) in 1882 called attention to the depletion of hepatic sly

glycogen fatty changes appear in the liver after exposure to these hepatotoxic agents Rosenfeld (39) observed that animals fed carbohydrate are, in general less susceptible to any drug which produces accumulations of fat in the liver Further more, after such poisonings the feeding of dextrose aids recovery of the animal Since the early reports of Whipple and Sperry (40), Opie and Alford (41), and Graham (42) on the resistance to chloroform or phosphorus poisoning of animals fed large amounts of carbohydrate or animals with livers containing large stores of glycogen, there have been many similar observations (43). The protective action of a high carbohydrate intake has also been noted in the prevention of hepatte damage following experimental ligation of the common bile duct (44), operation for Eck fistula (45), partial hepatectomy (46), and experimental poisoning with the mushroom Amantia bhalloides (47).

The various demonstrations of the lifesaving action of high carbohydrate intake on animals with experimentally damaged livers have been paralleled by clinical explorations of the therapeutic and prophylactic possibilities of administration of the contraction o

large amounts of carbohydrate. The recent experimental results outding up a man and his co workers have emphasized the therapeutic possibilities when account of the manufacture of the

stuffs on liver disease, it is gen here has been some work which

purports to show that high protein diets are as good or better than high carbohy drate diets. There is good evidence that in certain specific types of poisoning, namely those due to selentium (53) and arsphenamine (54), protein is definitely superior to carbohydrate in protective value. Indeed, in the exceptional case of sodium cyanide poisoning, high fat intake is better than either protein or carbohydrate (55). However, the evidence upon which the general superiority of protein is claimed is open to serious question. An examination of the data of Rav din and his co-workers (56–57) reveals that most of their companisons were made

asıs of com

parison is, of course, adequate protein-right calbonyunate. The gh proteinlow-carbohydrate Table 38 summarizes such a comparison (made in the authors' laboratory) for carbon tetrachloride poisoning in rats. It may be seen that the adequate protein-high carbohydrate due was definitely superior in lifesaving effect to both the high fat and the high protein diets. Chemical examination showed a correspondingly higher glycogen content of those poisoned animals which had been on the high carbohydrate thet.

It seems far to conclude that except in those instances where protein seems to text a specific action, its value depends upon its glycogeme and lipotropic proper test llence an adequate protein—high-carbohydrate diet is generally applicable In using such a diet, the uncreased requirements for the vitaniums of the B complex abould be satisfied. And in this connection it is important to note that large amounts of carbohydrate, together with a high dosage of thismine, would tend to produc fatty livers (58) unless counterbalanced by an adequate intake of choline or bustones amino acids.

Even this hnef survey of the subject points up the incompleteness of our present knowledge, especially as regards the particular effects of the various towns en

TABLE 38

SUPERIORITY OF HIGH CARBOHYDRATE ADEQUATE PROTEIN DIET
IN THE TRACTMENT OF CARBON TETRACHORIDE FOISONING
(MATTAR AND TAUBERHAUS [141])

Tive of Dirr	No DE		LIVER (All Values in Gm per Cent)			
	Rare		Glycogra	Fat	Protein A troops	
H sh fat H sh protein High carbohydrate	12 12 12	13 17 28	0 53 1 67 2 27	10, 85 5 68 5 06	3 50 3 34 3 66	

countered clinically. A systematic study of these and of the specific dietary combinations which are most effective in each case is certainly in order (59–60).

PHYSIOLOGIC BASIS OF INTRAVENOUS DEXTROSE TRERAPY FOR DISEASES OF THE LIVER

On the basis of Rosenbaum's observations and Rosenfeld's theories. Beddard (io) had suggested as early as 1908 that deterose be used clinically in large quanties to restore the depleted reserves of hepatic glycogen in cases of delayed poison by alter chloroform anesthesia. In addition to the administration of detrices by month and by rectal enemas. Beddard advised the intra-enous see of a 6 per cent solution. It is only recently, however, that the general introduction of adequate detrices therapy for hepatic disease has been shown to produce a definite decrease in mortality. In a cenes of cases in which acute hepatic insufficiency was treated with a ring and of carbohy drate given by mouth and intravenously. Jones (5) found that in a group of 10 cases observed from 1922 to 1925, in which 4:6-

patients were given a diet low in fat and supposedly high in carbohydrate, the mortality was 90 per cent. In only two instances was dectrose administered intra venously. In the next five years, with diets somewhat higher in carbohydrates (300–400 gm. daily) but with intravenous administration of dextrose in only four instances, there was 100 per cent mortality in 14 cases. However, in the years 930–35 when dextrose therapy was vigorous, 37 patients were treated with destin containing 400–500 gm. of carbohydrate daily, 26 of them receiving dextrose in travenously, and the mortality was lowered to 65 per cent. This author concluded "The more intensive the glucose therapy, the better the prognosis."

Despite these empiric observations, some difference of opinion still exists concerring the advantages of intravenous administration of dextrose if the patient can
take the necessary destrose or carbohydrate by mouth (62) But it should be
pointed out that the necessary amount of carbohydrate is supplied by the amount
of dextrose sufficient to raise the blood sugar to a level which will suppress the
output of hepatic sugar. Whereas the normal liver will respond to the usual post
prandial hyperglycetia, the "irritable" liver in acute toxemia may require a much
higher concentration of blood sugar to inhibit the formation of hepatic sugar.
That this is so was seen in the experiments previously described (p 270) in which
there was a prompt response of the acutely posoned liver in curtailing its output
of sugar when large doses of dextrose were given intravenously, while amail doses
had little or no effect (20).

Furthermore as Cori and Cori (63) have pointed out concerning the normal liver, "the blood sugar concentration and not the amount of glucose administered must be regarded as important for the rate of glycogen deposition in the liver." Consequently, when an attempt is made to protect a damaged liver by means of deposition of glycogen therein, the blood sugar concentration may have to be raised to a level which it may not be possible to obtain by feeding carbohydrates. In such cases intravenous infusion of dextrose is essential. The fact that extreme hyperglycemia so produced may result in glycosuria should not deter one from such vigorous therapy. As a matter of fact, this treatment has been successfully applied in diabetic patients with manifest or suspected liver disease (64)

Because of the glycosuna which may result from intravenous dectrose therapy, some physicians favor the routine use of insulin with the sugar. However, it should be pointed out that, unless the patient is diabetic, the indiscriminate use of insulin may defeat the very purpose for which the dectrose is administered. We have already referred to the evidence that, in the presence of sufficient insulin to maintain a normal constant blood sugar level, no additional insulin is necessary to obtain a normal hepatic response to administered sugar (65). Hence, the injection of insulin into a non-diabetic person can produce no additional hepatic effect, all though it does cause increased storage of glycogen in the muscles. Bridge (66) has shown that the administration of a certain amount of sugar intravenously to nor

mal rabbits resulted in higher levels of liver glycogen when it was given by itself than when insulin was injected simultaneously. This occurred despite the fact that the insulin caused no lowering of the blood sugar level. When the proportions of admisstered sugar and insulin are such that a lowering of the blood sugar results heliner is actually stimulated to pour out more sugar and is deprived of glycogen rather than replenished with it. Soskim Allweiss and Mirsky (29) have shown that the use of insulin with destrose in the treatment of toric non-diabetic animals shortens like animals recycling destrose alone live longer.

After prolonged intravenous injections of dextrose designed to suppress the sign producing mechanism of the liver the organ requires an interval to recover from the inhibition of dextrose formation so that hypogly cernia may appear from it as abours after the cessation of the infusion (67). This should be anticipated and itiated with small doses of carbohydrate by mouth or intravenously if necessary.

CLINICAL KETOSIS

Table 39 lists the abnormal physiological states and the cluncal conditions in which ketons is encountered. It also indicates the particular causative factors in when the tons is encountered. It also indicates the particular causative factors in when the state of the previous discussion in chapter x the fundamental disturbance underlying all ketons is a relative or absolute lack of enablydrate in the liver leading to an excessive brankdown of fat. However the enditions leading to this fundamental disturbance can be divided into three sub Furpir according to the manner in which it is brought about namely (a) disturbances in God mitake (b) impairment of liver functions and (c) endocrine disorder. It will be noted that there are a number of question marks in the table These are applied to certain of the endocrine mechanisms to indicate not only our figurentary knowledge as to the way in which they operate but also our lack of complete assurance that they operate at all in a particular condition. With these reservations however. Table 39 completely relates chinical ketons with our previous physiological considerations. Certain key references to more detailed consideration of the several conditions are also included in the table.

Von Gerke's disease and diabetes mellitus require some add tional comment. The former is exceptional in that it is the only condition in which ketous is associated with large stores of glycopen in the liver. But this glycopen is not available for use as is also evident from the fact that there is a low blood sugar level. Table 30 the glycopen in on Gierke's disease was therefore labeled abnormal hardly it is more likely that the glycopen itself does not differ from that found in normal livers but that the hepatic easy me systems are abnormal with a consequent mability to mobilize the glycopen were absent. As regards diabetes mellitus will be noted that the factor of insulin lack is designated relative or absolute. This is because, unlike exceptional and the land of the land of the same as if the glycopen were absent.

TABLE 39

CAUSATVE PACTORS IN VANIOUS STATES OF REPOSES (SUSKIN AND LEVINE [197]

	Deby	+	+	+ +	+
	Alka	+ +			
	Female Ser- Hor mone Facess			+	
	Adrenal Corts cal Ex-			- +	
E [roz]	Ante- rior Pitur- tury Excess	~~		~+~ ~	
O LEVIN	Rela tive of Absor- lute English Leck			++	
XIN AN	Abbor- mal Oly- coges		+		
222 (202	Erress De- namd for Car- heby-		+	+	+
OF A P.T.	Day Inched Gly- goges- ess		++++	+ +	
STATES	Exces- nwa Gly- cogen-		++++	+ +	
AKIDDS	Defr- event Carbo- bydrate Intake	+++	+		
CLOKS IN V	References	28.E.2.	ESSE.	(84 85) (85) (85) (88) (88) (88)	(901)
CAUGATTA LACTORS SIN VARIOUS SINTES OF A FIOSIS (SOSKIN AND LEVINE [107]	Cl nical States	Starvation 1.0.8 fit diet 2.0.8 fit diet 2.0.8 fit diet 3.0.8 fit diet 4.0.8 fit	Fever and infectious diseases A Acethoras Hepatita and early carbosis Advanced circulatory failure Von Gierke's disease	Disorders	Violent extruse
	ĺ	Disturb apoes 10	-megmi Jasm	Endocrme	Ĭ

whether in human diabetes mellitus there is an actual deficiency of insulin or whether there is an excess of opposing endocrine factors. From the practical thera results viewpoint, this, of course, makes little difference, since in either case the idministration of exogenous insulin will temporarily restore the disturbed endorine balance.

Scondary effects of kelosis —It is not at all certain whether the occurrence of ketone bodies in the blood and urine is in itself harmful. The evidence as to the tonicity of acctoacetic acid is contradictory, to say the least (77) Be that as it may, it is clear that the appearance of the ketones in excess of the amounts which can be metabolized by the peripheral tissues sets into motion a vicious cycle with a number of harmful secondary effects. The fact that the ketones are organic acids accessitates their neutralization, by sodium to preserve the normal pHI range of the blood and to enable their excretion by the tadincy. The ketonum as therefore accompanied by a loss from the body of fixed base and water. Further loss of clorder results from the vomiting which often accompanies ketons All these factors lead to dehydration and hemoconcentration, which, together with the loss of salts, result in an impairment of kidney function. When this occurs the ability of the body on metabolize and otherwise deal with the ketoacids rapidly dimmshes, and there begins a shift in the pHI of the blood to an extent incompatible with consciousness and life.

The post mortem findings, in individuals in whom Letons was the predominating cause of death, support our analysis of the pathological physiology. There are no specific organic lessons to be found. There is a cerebral capillary dilatation, privacular edema and acute degenerative changes in the cells of various parts of the certial nervous system. The findings in other parts of the body are those which are also seen in acute exsanguinating hemorrhage and in congestive heart failure. In general, therefore, the tissue pathology might very well be accounted for by acidoss, dehydration, hemoconcentration and cerebral amona.

The treatment of ketorss—For purposes of treatment, another classification of interest of claused ketosis may be made—namely, dishetes meltitus, on the one hand, and all other conditions on the other hand Diabetes is the only condition in which the original disturbance is a refative or absolute fack of insulin, and in diabetes the most essential part of the treatment is the early, adoptice, and position of insulin. This treatment will, of course, be rendered more effective by the simultaneous administration of adequate amounts of carbohy drate, "aler, and salt But the need for the hormone is paramount.

"act, and salt. But the need for the hormone is paramount it is equally important to remember that in non-diabetic ketosis the administration." "dinal principle of the

er under conditions in ary to accomplish this

purpose in the diabetic organism. The non-diabetic organism already has an op-

tained until the simple clinical and laboratory evidences of ketosis, dehydration, hemoconcentration, and hypochloremia have been abolished

INSTILIN PERISTANCE

In a number of clinical conditions the response of a patient to a given dose of insulin is less than that obtained in a normal individual. Diabetic patients who were formerly well controlled by a small dose of insulin may, with the advent of one of those clinical states, be poorly controlled even with very large insulin dosages. This phenomenon has been commonly referred to as "insulin resistance".

It is difficult to define normal insulin sensitivity very exactly, and there is no general agreement as to just bow abnormal the response must be in extent and duration to be called "insulin resistance". Lawrence (89) has reserved the term for instances in which the ethology is unknown. Strouse and his co workers (90) in their recent review of the subject chose to restrict their definition to cases of known or unknown ethology in which, after 48 hours' observation, 200 or more units of insulin could be administered without an appreciable lowering of the blood sugar

The various disturbances which might diminish the normal action of insulin may be listed as follows

r Poor absorption from the subcutaneous tissues

mally to its endocrine regulators

5 Unusual antibody formation to insulin or to other proteins present in insulin preparations

Various chincal cases have been reported in the medical literature in which one or another of the above factors bave been supposed to operate. But there is little good evidence that the suspected factor was actually responsible, and our knowledge of mechanisms is incomplete and is derived partly from clinical observation and partly from animal experimentation.

Root and his co workers (91) followed insulin absorption from the subcutaneous tissue by preparing a compound of insulin with radioactive iodine. This compound did not differ from insulin in its physiological activity, and the quantity present in an area in which it had been injected could be estimated from the degree of radioactivity. They found that their insulin compound was absorbed much more slowly from the subcutaneous tissues of diabetic patients manifesting insulin resistance than from the skin of other diabetic patients. The absorptive factor in the insulin resistant cases was confirmed by the fact that they responded smartly to insulin administered by the intravenous route.

srum and insulin. This has usually been interpreted as indicating the presence of an anti-missilin factor in the blood of insulin resistant individuals. Such a substance in gift be an antibody of some sort, of the effect might be non-specific and be due to an abnormally rapid rate of destruction of the added insulin. The possibility of bormonal antagonists is supported by the experimental evidence discussed in chapter xii and by the climical observations of increased requirement for insulin by diabetic patients coincidentally with the onset of thyroid or pituitary manifestit ons. As regards the formation of antibodies to insulin such cases occur but are rice (44 95). However, it has been observed that the insulin requirement of diabetics is likely to increase during the course of any allergic manifestations even though the patient is not allergic to insulin riself. The reported cases of insulin resistance in which an insulin antagonist in the blood has apparently been demonstrated.

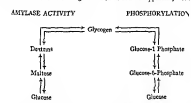


Fig. 74.—Intermediary substances: depending upon the mode of glycogenolys s. (Taubenhaus and Sokin [99])

itrated are not accompanied by the type of evidence which would permit a deter

Insul n resistance is most commonly encountered in infections and febrile states. The decreased effect of insulin has been variously ascribed to an overactivity of the terminal of the second of the

pos. fun

substantiated (21 29 96)

It was formerly thought that hepatic gly cogenolysis normally occurred through ambient that the place graded through determs to maistee and to flictose (Fig. 74). However Lee and Richter (70) who recently summarized the privious work on liver amylase and reported their own thorough studies on the salpest pointed out that (a) even the highest amylase activity found in the blood her and other organs is only of the order of 1/10 coo of the amylase activity of

PHYSIOLOGIC ACTION OF INSULIN IN SHOCK THERAPY OF THE PSYCHOSES

Schizophrenia and other psychoses have been treated with some degree of success by various forms of shock therapy including the induction of profound in sulin hypoglycenia. This influence of insulin has been attributed by some authors to a beneficial action of insulin upon the metabolism of the brain. This interpretation is not warranted.

The relationship between the blood sugar level and the utilization of carbohy drate by skeletal muscle was discussed in chapter xiv A similar relation between the blood sugar level and utilization of sugar has also been shown to hold for new c and brain tissues in dogs and in man (100 101 102) It will be recalled that the lower plateau in the S shaped curve which expresses the relation of the blood sugar level to utilization of sugar indicates that the latter cannot be depressed below a certain minimal rate by any degree of hypoglycemia (chap xiv p 151) Marked hypoglycemias may therefore drive the available supply of sugar from the blood below the minimal requirements of the tissues. Under such circumstances the muscle may have recourse to its stored glycogen or may perhaps turn to protein or fat as a source of energy. It is generally agreed however that nen e tissue has little stored carbohydrate and cannot utilize protein or fat. It follows that the nerve tissues during marked hypogli cemia are unable to maintain even the min mal rate of metabolism compatible with their well being. This explains the recent reports that prolonged insulin hypoglycemia has led to irreversible damage to the central nervous system in experimental animals (103) and to similar pathologic changes and mental deterioration in schizophrenic patients (104) It may be con cluded that insuling shock, therapy has been well named?

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CHAPTER XXIII

COMPARATIVE PHYSIOLOGY OF DIABETTS TT WAS fortunate for the development of the science of metabolism that I WAS lortunate for the development of the science of metabolism that Mering and Minkowski, in 1890, chose to depancreature dogs. In this species Menng and Minkowski, in 1890, chose to departreature dogs in this species and is characterized by hyperglycenia, in the partraine diabetes develops scately and is characterized by hyperglycenia, pancreatic diabetes develops scuted and is characterized by hypersylvemia, betonis etc. It bears a striking resemblance to gycosuria, polyuria, polydypsia, actosis etc. 11 bears a striking resemblance to dusters mellitus in man even though the syndromes diverge in several dedishetes mellitus in man even though the syndromes diverge in several de-tails The effect of Menng and Minkowski a work on the dog was to advance our tain. The effect of Menng and Minkowski a work on the dog was to advance our according to the effect of Menng and Minkowski a work on the dog was to advance our according to the effect of Menng and Minkowski a work of the effect of Menng and Menng and Minkowski a work of the effect of Menng and Menng ental preparation
As early as 1879. Langendorf (1) had removed the Pancreas of chickens and As early as 1879, Langendori (1) had removed the bancress of chickens and they apparently died specified birds did not exhibit glycosuria and they apparently died to the control of the c pigeons The operated birds and not exhibit glycosuria and they apparently died in extreme emacation consequent to a loss of appetite. The clear relationship be in extreme emacation consequent to a loss of appetite. in extreme emacuation consequent to a loss of appetite. The clear relationship of these paners also function and normal carbohydrate metabolism could not have ween pancerate function and normal cathonyurate metabolism could not nave (2) and the north with domesticated birds. In 1897 Minkowski (2) the defended from this north with domesticated birds in 1897 Minkowski (2) and the north with the north with domesticated birds. mental preparation nem areduced from this work with domesticated birds. In 1501 Minkowski (2)

ontimed the observations of Langingford and extended his studies on the effects

of monomials and are also as the contract of the construed the observations of Langendori and extended his studies on the enects of papers account to several other species. Since that time there have appeared to paper account to several other species. or panerestectomy to several other species once that time there have appeared sporadic studies in comparative disbetes, notably from the laboratories of types of the studies of the species of the speci 3).4.5.6), Lukens (7.8), Mirsky (9 to 11), and Houssay (12, 13)
Table 42 summarizes some of the characteristics of the syndromes which follows: grandor nuclea in comparative chareces, notating from the manufacture of the state Asone 42 summarues some of the characteristics of the syndromes which follows an apparent of the syndromes which the syndromes paintreasterctomy in the various species which have been studied. Diabetes meititude in man its included for comparison. It can be seen that the effects of the removal of the object of the removal of the object of the removal. in man is included for comparison. It can be seen that the cliects of the removal of the grand upon blood sugar, sugar exerction, protein breakdown, kelosis, and n me giand upon blood sugar, sugar exerction, protein breshdown, ketosis, and time of survival vary wilely but not in any obviously related fashion and the of survival vary wilely but not in any obviously related fashion and the obvious a time of survival vary widely but not in any obviously related tashion. I have in both the dog and the cat the diabetic state is severe, as judged by Silversurfa. nontine dog and the cat the diabetic state is severe, as junged by \$10 courts and the protein breakdown, but while belows in the cat is very severe it is generally mide. puren neestdown, but while ketons in the cat to very severe it is generally mild hyper in the dog. The depancers turned page and goar, on the other hand, exhibit mild hyper and the dog to the depancers turned page and goar, on the other hand, exhibit mild hyper and the dogs are the dogs and the dogs and the dogs are ne include the department and place and four the other hand, empire much have the service of protein breakdown above the normal and gly counts, with lettle, if any, increase of protein breakdown above the normal and gly counts, with lettle, if any, increase of protein breakdown above. gyremus and glycosuria, with httle, if any, increase of protein oreascoorn asove the normal rate. The goat has a correspondingly mild ketonuria. the normal rate. The good has a correspondency mild ketonura, but the diabetic figure normal rate. The good has a correspondency mild ketonura, which, however, pig develops a very high level of ketone boddes in the blood. And any a shadow of develops a very high level of ketone boddes in the blood. ing generous a very high level of ketone bodies in the billood which, however, does not seem to extrt any effect on the acid base balance. The duck and chicken does not seem to ever any effect on the and base balance. The dock and entering only occasionally and transcently. .44 section to counts after panertatectomy unit sections that produces Hovever, removal of the gland in these hards produces movers, removal of the giant in toese thrus from specific the depart. The depart of the giant in the depart. El cosma pit no retorn and sact not suppost institutions institutions institutions institutions institutions institutions institutions in the company in the company institution in the company in the co The mechanisms responsible for these species diff

TABLE 42 Species Variation in the Effects of Pancheatectomy

	ENCHS	£ 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5		(28)
	XXXXXX	ment, 37 units per day ment 57 units per day ment 57 units per day ment 58 ago units per day ment 58 ago units per day y mompliets ment 1 transent ment 1 tran		
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SURVEYAR	(Dave)	10 5 3 38-120 41 163 41 163 41 163 6 6 6 6 6 100+ 100	tell tus	
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Urde Excertion (Gr/Eg/Day)	N trogen	0 00 04 		o 13-0 35
5	Sugar	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	(0 77 0
Secar Cort)	D abet e	341 300-700 310-345 310-345 310-345 400-700 100-312 30-13 110-30 310-310 110-30 10-30		200-600
BLOOD SPEAR (Ma FER CENT)	No mad	(60-90 60-100 30-70 30-710 30-		00-00
Sterns		film flooring of the control of the		Man

dated to date. However, there are indications that several different factors may using 10 using 2100NEVER there are indications that several different factors may play a role in modifying the diabetes of the various animals. On the hasis of the oppoint activities of insulin on the one hand and the hormones of the anterior posms activities of insume on the one mand and the normanes of the anterior hypophysis the adrenal cortex and the thyroid on the other it might be suppygognysos one autenas correct musu one cosystem was the metace of mugat toe sup-posed that the mild disabetes of some species may result from a characteristic or in bred wakeness of the endocrine opponents of the pancreas. This seems to be trie or the pig which exhibits a diabetes similar in its characteristics to that of the hypothysectomized-depancreatized or adrenalectomized-depancreatized dog or appropriate administration of antenor pituitary extracts intensifies the diabette state of the p g (7). However, the hypothesis of variable endocrine halance does not ac on the $p \in U$) answerer the hypothesis on variable characterise maintenance ones not account for the modification of diabetes seen in other species. Thus, pituitary hor

wome for the monuneation of disocres seem in other species 3, 2103, pive mores do not induce manifest diabetes in the departers tized duck (9) ω_{mea} wo not immore mannest charactes in the department duck (9). Species differences after panereatectomy might also be due to variations in the relative importance of factors other than insulin removed with the pancreas—for resulve unportance of factors other man mamma removes while the planetess—for example, the lipotropic function of panetestic secretions. This seems to be true in the case of monkeys Coll p (14) and Fulton (15) both reported that pancrate tomy us the monkey results in a mild diabetes resembling that of the Housesy dog Now, many results in a time tracers assumed that department monkeys on a diet supplemented with pancreatin found a severe d abetic state resembling that of the prometrical with panereating fourth a severe to asset it spatially that it is a severe to asset it is a severe to a severe to a severe to a severe to that the absence of hipotropic factors could not explain the failure of the duck to de dop dubetes. The inclusion of pancreatin in the diet of depancreatized ducks. priorp materia. The increasion of paintream in the mense neight loss but did not led to hypergly centra or glycosura (9) The a cells of the islands of Langerhans which in panetratectomy are removed along with the insulin producing S.c.ells any also play an as yet undiscovered role in influencing the diabette syndrome may also play an as yet undiscovered role in inducencing the dissociated systematics. Allocanized degs in which the a-cells are undisturbed exhibit more intense given suna less ketosis and longer surrival without insulin treatment than do de

H may well be that the apparent variation in the diabetic syndrome of a par at may well be that the apparent variation in the charge symmetric of a plant to the mean plant removal of insulin producing issue that the apparent variation in the charge of the mean plant removal of insulin producing issue that the mean plant removal of insulin producing issue that the apparent variation is the charge of the mean plant removal of insulin producing issue that the apparent variation in the charge of the mean plant removal of insulin producing issue. remaraperers may be due to the incomplete removal of insum producing issue performed rabbits have a prolonged survival time and little ketosis (r6). panereatized dogs (19) New Control of the C

This a severe autoxanized rabouts (in a nien presumacity au presis are destroyed).

This a severe acidosis with a ketonemia of 120 mg per cent (17) Alloxanized rats (28).

On the bass of his observations Minkowski (2) made the generalization that (18) show no striking differences from departerealized rats charged animals of the observations alinkowski (2) made the generalization that the charged animals suffer from a more intense pancrestic diabetes than do denivorous animals suffer from a more intense painteaue discussion and the Herbivora. He and Weintraud (20) showed that unlike chickens p. geometric discussions and denivorous animals suffer from a more intense passes of the property of t are necroscora. He and Weintraud (20) showed that unine chieckins if Front and ducks the carm orous owls and hawks exhibit immediate fi) cosuma after the carm orous owls and hawks exhibit immediate for course for the course of the course and unces the carmerous owns and haves eminer minimum by communa and programme and programme and extended the work on owns and programme and progr also attempted to change the response of the duck by a prelumnary period of meat feeding No conclusive results were obtained, although some of the meat feed ducks did develop a certain degree of hyperglycemia and glycosum. The previous die tary habits of an animal might influence the characteristics of its pancreatic dia betes by affecting the Secretory activity of certain endocrine glands or by setting the metabolic reactions in the liver in one or another direction. In the latter connection it might be well to recall the observation of F. G. Young (21) that the feeding of meat or non protein extracts of meat increases the severity of ketosis in dogs with metahypophyseal diabetes.

Whatever the causes of species difference in diabetes may prove to be, the subject is by no means one of academic interest only. It has already been pointed out that the etiology of diabetes mellitus in the human is unknown and that in the majority of cases it is evidently not due to pancreate pathology (chap xxii, p 465). It may well be that further and more exact knowledge of the causes of species variation in the diabetic syndrome could suggest possible etiologic factors in man. For this purpose, further work comparing alloxanized animals and studies on the gluconcogenetic response of various species to phlorhizm should be profitable.

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CHAPTER XXIV

PRESENT FRONTIERS OF RESEARCH IN METABOLISM

ALTHOUGH this volume has dealt primarily with the metabolism of car bohydrate, it has been necessary to consider the metabolism of protein and fat to a considerable extent. As a matter of fact, the division of the subject of metabolism into three compartments, related to the three major food stuffs, is largely artificial, depending upon the limitations of the authors rather than upon any real separation of the subject matter. In the light of more recent knowledge of intermediary metabolism, it seems likely that we shall soon cease to distinguish between the metabolisms of the different foodstuffs, once they have gone beyond certain stages, for, eventually, all of them give use to very similar intermediary products, namely, the α and β ketoacids

Interrelationships between carbohydrate protein, and fat metabolish

Figure 18 (p 54) presents a composite scheme of the main pathways connecting the metabolism of carbohy drate, protein, and fat The supporting evidence is drawn from in 110, periuson, and in vitro experiments on different animals and under different conditions. No single animal, organ, or type of tissue has been shown to be capable of performing all the reactions in the scheme Indeed, there is evidence that certain tissues lack the ability to carry on many of them. The scheme therefore applies to the organism as a whole, i.e., a certain tissue may carry the degradation of a foodstuff or the synthesis of an intermediate product to a given point and then pass on its end product, by way of the blood, to another tissue which completes the process.

If the tentative scheme shown in Figure 18 is substantiated by future work, it will be possible to speak of a "final common pathway" for all the foodstulfs. The intermediary metabolities composing the tricarborytic acid cycle (see chap in) will then be regarded as a metabolic pool to which all the foodstulfs contribute and from which they can be regenerated (amination, CO, firation). This will obviate much of the former discussion as to the interconvertibility of a particular foodstuff into another, for it will be realized that none of them are interconvertible in the sense that the constituent atoms of one pass directly and in a body into the other, while all of them are interconvertible, in the sense that the augmentation of the pool by a certain amount of intermediary material derived from any food

stuff may displace an equivalent amount of intermediary substance from the pool for the synthesis of another foodstuff

It might be objected that if the interchangeability of foodstuffs were as complete as is indicated by the scheme it should be possible to maintain adequate nu inton on a dict composed solely of any one of the foodstuffs. But we know that only protein—and indeed only certain proteins—can be used in this way, and then for limited periods of time only. The answer to this objection lies not in any lack of interconvertibility but in the fact that animal roetabolism is incomplete Animals cannot synthesize certain essential food materials but must obtain them from plant and mineral sources. These essential accessory food factors comprise (i) the essential amino acids. (a) the essential fatty acids. (3) the vitamins and (4) the minerals. It is happens that only a mixed dictary of natural foods will contimum the necessary amounts of all the accessory food factors of all the accessory food factors.

SIGNIFICANCE OF 18 Prire RESULTS

The best available scheme for the dynamics of carbohy drate metabolism was presented in chapter in But it must be emphasized that despite its general plausibility and inner logic it is only a tentative outline. The data for it are derived from work done with intact with eviscerated and with hepatectomized animals aid from observations made after the removal of various endocrine glands etc. The proparations used for in tifro work include organ slices miniced tissues and thryme extracts.

The various techniques of in vitro work have been invaluable for the development of our present concepts of intermediary metabolism but they suffer from several inherent limitations which are not always appreciated or emphasized. Even itssue sices in which there is presumably the least physical damage to in dividual cells do not exhibit quite the same metabolic behavior as do the parent bissues in time. For example, liver shees cannot be induced to deposit glycogen (except rarely and to an insignificant degree) (i = 3) as the organ so readily does in time. The liver since in time appears to be exclusively in the phase of glycogenolysis. In this connection it may be pertinent to consider the fact that the inlate liver possesses a dual blood supply each supply differing in rate of flow presure and oxygen and CO, contents The cells of the liver slice in vitro must function in a uniform medium. Turning from liver to brain we note that the high test is not oxygen consumption of cortical slices is only from one third to one half the oxygen consumption of whole brain in vitro (4, 5, 6). Obviously, some unknown factors modify metabolism when tissues are separated from their normal en alternative.

Mincing of tissues introduces even more serious deviations. For example, while an intact thin muscle (diaphragm or abdominal muscle) retains its ability to deposit glycogen from glucose (7 8 9) and can also use the glucose in the medium

CHAPTER XXIV

PRESENT FRONTIERS OF RESEARCH IN METABOLISM

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INTERRELATIONSHIPS BETWEEN CARBOHYDRATE

Figure 18 (p. 54) presents a composite scheme of the main pathways connectrigure to W 34) presents a composite scheme of the main paramaps connects ing the metabolism of carbohydrate, protein, and fat The supporting evidence is ang the metabolism of carbonyunare, process, and he are supporting evidence as drawn from in 1900, perfusion, and in 1910 experiments on different animals and under different conditions. No single animal, organ, or type of fissue has been shown to be capable of performing all the reactions in the scheme Indeed, there stown to ne capanic of performing on the reactivity at the statement threety, three is evidence that certain fissues lack the ability to carry on many of them. The as evadence that betteam thousand start the animaly to tailiy on many or them. The scheme therefore applies to the organism as a whole, i.e., a certain tissue may carry the degradation of a foodstuff or the synthesis of an intermediate product carry the degeneration of a societies of the synthesis of the intersections product, hy may of the blood, to another

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suff may displace an equivalent amount of intermediary substance from the pool for the synthesis of another foodstuff

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SIGNIFICANCE OF 18 Citro RESILIES

The best available scheme for the dynamics of carbohy drate metabolism was presented in chapter in But it must be emphasized that, despite its general plausibility and niner logic, it is only a tentative outline. The data for it are derived from work done with intact, with eviscerated, and with hepatectomized animals and from observations made after the removal of various endocrine glands etc. The preparations used for in ritro work include organ slices, minord tissues, and mymetritures.

The various techniques of an extre work have been invaluable for the development of our present concepts of intermediary metabolism but they suffer from several mherent limitations which are not always appreciated or emphasized Even tissue slices, in which there is presumably the least physical damage to in dividual cells do not exhibit quite the same metabolic behavior as do the parent tissies in tro For example, hver shees cannot be induced to deposit glycogen (except rarely and to an insignificant degree) (1, 2, 3), as the organ so readily does in two The liver shee in trito appears to be exclusively in the phase of gly cogenolysis. In this connection it may be pertment to consider the fact that the intact liver possesses a dual blood supply, each supply differing in rate of flow, pressure, and oxygen and CO, contents The cells of the liver slice in vitro must function to a uniform medium. Turning from liver to hrain, we note that the high est in ribo oxygen consumption of cortical slices is only from one-third to one half the crygen consumption of whole brain in ere (4 5,6) Obviously, some unknown factors modify metabolism when tissues are separated from their normal en vironments.

Mincing of tissues introduces even more serious deviations. For example, while an intact thin muscle (disphragm or abdominal muscle) retains its ability to deposit glycogen from glucose (7, 8, 9) and can also use the glucose in the medium

for energy purposes, mincing interferes with the entry of glucose into the cells for either purpose

Cell free extracts are a step further removed from normal relationships. The generally used muscle extract of Meyerhof (10) contains the stable systems soluble in 0.6 per cent potassium chloride or water. The water insoluble enzyme proteins, such as myosin, are not present, and the creatine phosphate hydrolyzed during the preparation of the extract. It is obvious that the carbohydrate metabolism of such an extract is quantitatively and qualitatively different from that of intact muscle. For example, it is well known that many tissues (e.g., muscle) show a greater breakdown of carbohydrate and a larger formation of lactic acid during anoxia than during adequate oxygenation. The inhibitory influence of oxygen on the rate of glycolysis is known as the "Pasteur effect" (i.i., i.i). The exact mechanism of this effect in intact tissues is not entirely clear. Among the factors which may be involved are (e.g. the hreakdown of organic phosphate during anoxia, providing excess inorganic phosphate, which would orient the reactions toward glycolysis (i.j.), and (b) the fact that many enzyme proteins involved in glycolysis are active in the -SH state (reduced) and may therefore be inhibited by an in creased oxygen tension (i.4, i.5). Whatever its mechanism, the Pasteur effect is an important regulatory phenomenon in carbohydrate metabolism in vivo—a mechanism which is completely lacking in tissue extracts.

From even these few considerations it becomes obvious that extreme caution is necessary in applying in vitro data to the elucation of in vivo metabolism. Fur thermore, a homogeneous cell free enzyme extract, even if it contained all the cell proteins in their in vivo proportions, would not be very comparable to the living cell. In the latter, heterogeneity and structural separation, etc., make it possible to have a number of zones within a single cell, each differing as to pH, mineral composition, etc., and each varying in metabolic activity. External influences, both physical and chemical, may therefore influence the metabolism of the cell by inducing changes in its internal structure. For example, the structural change induced in myosin by the nerve implies activates carbodydrate breakdown and alters the rate of oxygen consumption. The rate of metabolism is also influenced in unknown fashion by thyroid hormone or by dinitrophenol. These substances may act by bringing together links in the respiratory chain which, although al ways present in the cell, are usually separated from each other in some way

More specifically, Stannard (16, 17), Korr (18), and others (19, 20) have shown that, in certain tissues, work or chemical stimulation not only raises the rate of oxygen consumption but alters the pathway by which it is used The low oxygen consumption of these tissues at rest is resistant to the influence of cyanide despite the presence of the cytochrome system on which the poison acts. When the tissues are stimulated, the oxygen consumption rises and becomes sensitive to cyanide

Apparently the stimulus in some way links the idle cytochrome system to the de Apparently the stimulus in some way image the rule cytochronic system to the de bydrogenases Similarly, the work of Sacks (21, 22, 23) and of Flock and Bollman nydrogenases Summarry, the work of Sachs (21, 22, 23) and of Flock and Bournan (42, 23) understees that the scheme of phosphorylations via adenosme triphosphate (42.23) moreates that the scheme of phosphorylations via adenosme triphosphate (ATP), outlined in chapter IV, may apply to muscle at rest but may not be wholly (ALL), outlined in chapter IV, may apply to muscle at rest but may not be wholly valid for the same tissue during work. Although this work has been criticated Yama or the same Ussue during work Although this work has been counted (26, 27), it should put us on guard against regarding the presently accepted meta

_{oue} scnemes as either complete or unai In addition, it should again he recalled that the scheme of intermediary carbo in addition, it should again be recaused that the scheme of intermentary CHIDO by draft metabolism has been constructed from data obtained in different animals. bolic schemes as either complete or final nywest incremental has need constructed from data obtained in unierent animals and issues. It is a composite picture, and not every tissue or organ conforms to it Thus the liver produces very little lactic acid, and yet it can build up hexoses and when were produces very intre facts acid, and yet it can build up nexoses and glycogen from lactate (48). For the liver therefore the scheme requires modificate. eyougen non uscrate (28) For the uver therefore the scheme requires modules too to account for these phenomena. To eite another example, skeletal muscle tion to account for these phenomena. To eite another example, secretar innertent tusive requires insulin for good rates of glycogen synthesis from glucose (7, 29). The Library requires insulin for good rates of grycogen synthesis from gricose (1, 29). Life hart and kidney, on the other hand, may deposit greater than normal amounts neart and Ridney, on the other hand, may deposit greater than normal armon of gyrogen when the blood sugar level is high but insulin is absent (30, 31)

by $\omega_{S^{(1)}}$ when the mood sugar level is high nut insum is ansent (30-34). Taking into account all the pitfalls inherent in the various m silvo technics, we may sum up by stating that, when a reaction or a series of reactions is shown to may some up stating that, when a reaction or a series of reactions is sometime.

The series of reactions are sometimes as some reactions can, but do not neces the series of the series suly, occur in the living infact organism. A negative is tife result is wholly me. conclusive, since it may simply depend upon the conditions of the experiment. All concursive, since it may simply depend upon the conditions of the experiment, in wito data must eventually be checked in sito, in order to acquire serious significant to the conditions of the experiment of the checked in sito, in order to acquire serious significant to the checked in sito, and the checke on the data must eventually be enecked in sits, in order to acquire serious as moderne in our concepts of normal metabolism. For this purpose, the labeled nour concepts of normal metabonsm for this purpose, the ascient account of the concepts of normal metabonsm for this purpose, the ascient of the concepts of normal metabonsm for the concepts of no 33) The intravital staming technique of Gomori and others (34-35) and spectro 33) the intravital staining technique of Comori and offices (34, 35) and the photometry of living tissues (36, 37, 38) also hold promise for the future

The previous discussions concerning the action of insulin (chap xvi p 180) the previous discussions concerning the action of insulin (cnap xvi p 100) and the telear that glycogen deposition from glucose could proceed at a relatively and it clear that glycogen deposition from glucose could proceed at a renatively slow rate in the complete absence of the hormone. Apparently, the enzyme sysour rate in the complete absence of the hormone. Apparently the enzyme systems necessary for the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in the completely defined by the polymetrzation of glucose are present in cons necessary for the polymentation of glucose are present in the coinqueries we hanced by more than a three properties of their activity can be markedly enhanced by more than the rates of their activity can be markedly enhanced by pointreatized animal, but the rates of their activity can be markenly eminance up to the form one is, therefore not a necessary cell enzyme itself but a republic of the proposed animals. lator of rates of reaction. This point of view is supported by a consideration of the MOUT OF TRIES OF reaction. This point of view is supported by a consuctiation of the amount of insulin which must be administered to restore the normal metabolic amount of insulin which must be administered to restore the normal insulations for its departmental animal. This has been shown to be of the order of 2507 eastern a departeratized animal. This has been shown to be or the order to gave for a toky dog per day. Even if no destruction of the administered insulin or more to gave the same of the administered insuling the same of t are a to-kg dog per day Even if no destruction of the administered insuin or cured, this would result in a concentration of approximately 0 37 per 100 gm of curred, this would result in a concentration of approximately 0.37 per 100 gm to tussee water. This order of magnitude is far lower than that of the concentration.

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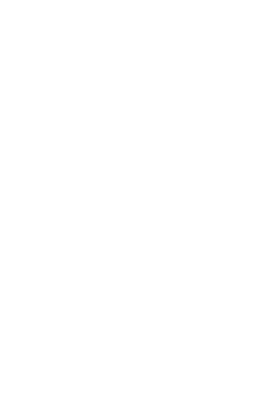
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